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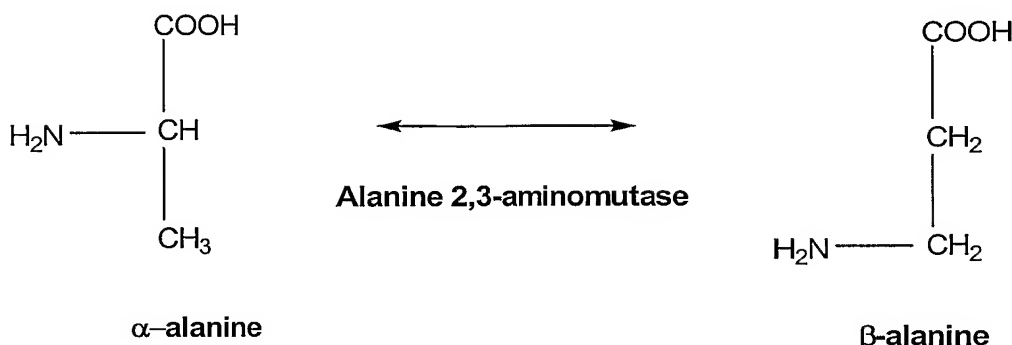
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(54) Title: IMPROVED ALANINE 2,3-AMINOMUTASES AND RELATED POLYNUCLEOTIDES



(57) **Abstract:** The present invention is directed to polypeptides that have enhanced alanine 2,3-aminomutase (AAM) activity and/or thermostability relative to the wild-type enzymes that have incidental AAM activity as a result of cross reactivity with alanine. In addition, the present invention is directed to a polynucleotides that encodes for the AAM polypeptides of the present invention, to nucleic acid sequences comprising the polynucleotides, to expression vectors comprising the polynucleotides operatively linked to a promoter, to host cells transformed to express the AAM polypeptides, and to a method for producing the AAM polypeptides of the present invention.

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IMPROVED ALANINE 2,3-AMINOMUTASES  
AND RELATED POLYNUCLEOTIDES

FIELD OF THE INVENTION

[01] The present invention is related to the field of enzymology, and particularly to the field of alanine 2,3-aminomutase (AAM) enzymology. More specifically, the present invention is directed to alanine 2,3-aminomutase polypeptides having improved enzymatic activity (*i.e.*, high substrate turnover) and stability, and to polynucleotides sequences encoding for the improved alanine 2,3-aminomutase polypeptides. The present invention is useful because the alanine 2,3-aminomutase polypeptides can be coupled to other enzymes to produce synthetic organic chemicals, such as pantothenic acid or 3-hydroxypropionic acid in high yields.

BACKGROUND OF THE INVENTION

[02] Organic chemicals such as organic acids, esters, and polyols can be used to synthesize plastic materials and other products. To meet the increasing demand for organic chemicals, more efficient and cost-effective production methods are being developed which utilize raw materials based on carbohydrates rather than hydrocarbons. For example, certain bacteria have been used to produce large quantities of lactic acid used in the production of polylactic acid.

[03] 3-hydroxypropionic acid (3-HP) is an organic acid. Several chemical synthesis routes have been described to produce 3-HP, and biocatalytic routes have also been disclosed (WO 01/16346 to Suthers et al.). 3-HP has utility for specialty synthesis and can be converted to commercially important intermediates by known methods in the chemical industry, *e.g.*, acrylic acid by dehydration, malonic acid by oxidation, esters by esterification reactions with alcohols, and 1,3-propanediol by reduction.

[04] The compound 3-HP can be produced biocatalytically from PEP or pyruvate, through a key beta-alanine intermediate (FIG. 1). Beta-alanine can be synthesized in

cells from carnosine, beta-alanyl arginine, beta-alanyl lysine, uracil via 5,6-dihydrouracil and N-carbamoyl-beta-alanine, N-acetyl-beta-alanine, anserine, or aspartate. However, these routes are commercially unviable because they require rare precursors or starting compounds that are more valuable than 3-HP. Therefore, production of 3-HP using biocatalytic routes would be more efficient if alpha-alanine could be converted to beta-alanine directly (FIG. 1). Unfortunately, a naturally occurring enzyme that inter-converts alpha-alanine to beta-alanine has not yet been identified. It would be advantageous if enzymatic activities that carry out the conversion of alpha-alanine to beta-alanine were identified, such as an alanine 2,3-aminomutase. Accordingly, it is one object of the present invention to identify enzymes with improved alanine 2,3-aminomutase activity.

[05] Lysine 2,3-aminomutase (KAM), which catalyzes the anaerobic interconversion of lysine to beta-lysine, was first described by Barker in *Clostridium* SB4 (now *C. subterminale*) catalyzing the first step in the fermentation of lysine. KAM has been purified from *C. subterminale*, the gene cloned and expressed in *E. coli*. See e.g., U.S. Pat. 6,248,874, which issued on June 19, 2001 to Frey *et al.*, the whole of which is hereby incorporated herein by reference. The specific activity of purified KAM from *C. subterminale* SB4 cells has been reported as 30-40 units/mg (Lieder et. al., Biochemistry 37:2578 (1998)), where a unit is defined as  $\mu$ moles lysine/min. The corresponding purified recombinantly produced KAM had equivalent enzyme activity ( $34.5 \pm 1.6 \mu$ moles lysine/min/mg protein). See U.S. Patent Application Publication No. 2003/0113882 A1, which published on June 19, 2003 to Frey *et al.*, the whole of which is incorporated herein by reference.

[06] Based upon the sequence of the KAM from *C. subterminale*, KAM genes have been annotated in the genomes of other organisms. However, in most cases, the enzymatic activities of the polypeptides encoded by these genes have not been confirmed. Exceptions are the *B. subtilis* gene (Chen, D., Ruzicka, F.J., and Frey, P.A. (2000) Biochem. J. 348:539-549), and the *Porphyromonas gingivalis* and *F. nucleatum* genes. The *B. subtilis* KAM, encoded by the *yodO* gene, is more resistant to  $O_2$  than the *C. subterminale* KAM, but it is markedly less active. As reported by Frey, the *B. subtilis* KAM has a specific activity of only 0.62 U/mg.

[07] *C. subterminale* SB4 KAM has been reported to have some cross-reactivity with L-alanine, converting it into beta-alanine. See U.S. Patent Application Publication No. 2003/0113882 A1. WO 03/062173 and WO 02/42418 disclose the first reports of AAM activity based upon modification of *kam* genes. In these applications, the synthetic *aam* genes had AAM activity as detected by the complementation of a  $\Delta$ panD *E. coli* strain. However, because alanine is not the natural substrate for this enzyme, the activity for this conversion is substantially less than the activity for conversion of lysine — its natural substrate. The AAM activity of a variant of *B. subtilis* KAM that also had AAM activity at approximately 0.001 U/mg. It is an object of the present invention to provide polynucleotides encoding a polypeptide having substantially enhanced AAM activity over that found in the wild-type enzymes.

## SUMMARY OF THE INVENTION

[08] The present invention has multiple aspects. In one aspect, the present invention is directed to polypeptides that catalyze the reaction of FIG. 1. In one embodiment of this first aspect, the present invention is directed to a polypeptide having alanine 2,3-aminomutase (AAM) activity, preferably as measured by the assay of Example 8, and,

- (a) having a polypeptide selected from the group consisting of SEQ ID NO: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36, 38, 40, 42, 44, 46, 48 and 51;
- (b) having an amino acid sequence which has at least 98% homology, with the amino acid sequence selected from the group consisting of SEQ ID NO: 2, 22, 28, 32, and 36;
- (c) having an amino acid sequence which has at least 99% homology, with the amino acid sequence selected from the group consisting of SEQ ID NO: 4, 6, 8, 12, 16, 24, 26, 30, 34 and 40;
- (d) being a polypeptide encoded by a nucleic acid sequence which hybridizes under high stringency conditions with either (i) the nucleotide sequence of SEQ ID NO: 1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25, 27, 29, 31, 33, 35, 37, 41, 43, 45, 47 or 49; (ii) a subsequence of (i) of at least 100 nucleotides, or (iii) a complementary strand of (i) or (ii) (J. Sambrook, E. F. Fritsch, and T. Maniatis, 1989, Molecular Cloning, A Laboratory Manual, 2d edition, Cold Spring Harbor, N.Y.); or
- (e) being a variant of the polypeptide of (c) comprising a substitution, deletion, and/or insertion of one to six amino acids therefrom and having AAM activity from about 1 to about 30  $\mu$ M  $\beta$ -alanine produced /hour 1 cell OD at pH 7.0-7.6, 25°C.

[09] Collectively, the polypeptides of (b) and (c) above are referred to herein as "homologous polypeptides." For purposes of the present invention, the degree of homology between two amino acid sequences is expressed as "percent homology," "percent identity," "% identity," "percent identical," and "% identical" are used interchangeably herein to refer to the percent amino acid sequence identity that is obtained by ClustalW analysis (version W 1.8 available from European Bioinformatics Institute, Cambridge, UK), counting the number of identical matches in the alignment and dividing such number of identical matches by the length of the

reference sequence, and using the following default ClustalW parameters to achieve slow/accurate pairwise optimal alignments – Gap Open Penalty:10; Gap Extension Penalty:0.10; Protein weight matrix: Gonnet series; DNA weight matrix: IUB; Toggle Slow/Fast pairwise alignments = SLOW or FULL Alignment.

[10] In one embodiment, the present invention is also directed to an AAM polypeptide as described herein in isolated and purified form.

[11] In another embodiment, the present invention is directed to an AAM polypeptide as described herein in lyophilized form.

[12] In yet another embodiment, the present invention is directed to a composition comprising an AAM polypeptide as described herein and a suitable carrier, typically a buffer solution, more typically an aqueous buffer solution having a pH between 6.0 and 8.0. The composition may also be in a lyophilized form.

[13] The novel AAM polypeptides of the present invention have significantly enhanced AAM activity relative to the wild-type KAM polypeptides from which they are ultimately derived. By significantly enhanced AAM activity is meant that the AAM polypeptide of the present invention has an AAM activity within the range of about 1 to about 32  $\mu$ M  $\beta$ -alanine produced/hour 1 cell OD (units), preferably from about 10 to about 32 units, more preferably from about 20 to about 32 units; most preferably from about 25 to about 32 units.

[14] Preferred AAM polypeptides of the present invention have an amino acid sequences of SEQ ID NOs: 2, 6, 12, 16, 20, 24, 28, 30, 32, 34, 38, 44, 46 or 48; more preferably they have an amino acid sequence of SEQ ID NOs: 6, 12, 28, 34, 46 or 48; most preferably, they have an amino acid sequence of SEQ ID NOs: 28 or 34.

[15] One of the grandparent molecules is the KAM of *Bacillus subtilis*, which had no detectable AAM activity. The DNA encoding this grandparent molecule was modified as described in WO 03/062173, entitled “Alanine 2,3-aminomutase,” to produce a polypeptide having a detectable alanine 2,3-aminomutase activity.

[16] In the present application, the applicants utilized as one parent molecule a polynucleotide sequence of SEQ ID NO: 58, which encoded the 471 residue polypeptide of SEQ ID NO: 59 and which exhibited an AAM activity of

approximately .001 U/mg (units/ mg of cell mass). The molecule of SEQ ID NO: 59 differs from the wild-type *B. subtilis* KAM, which had no detectable AAM activity, by having the following four (4) amino acid substitutions: L103M, M136V, Y140H and D339H.

[17] In yet another embodiment, the present invention is directed to a polypeptide having from about 1 to about 32 units of AAM activity and typically varying from the polypeptide of SEQ ID NO: 59 by 1-7 amino acid residues, more typically by 1-6 amino acid residues, even more typically by 1-5 amino acid residues, and most typically by 1-4 amino acid residues.

[18] In its second aspect, the present invention is directed to a polynucleotide sequence that encodes for the correspondingly referenced AAM polypeptide. Given the degeneracy of the genetic code, the present invention is also directed to any polynucleotide that encodes for the above referenced AAM polypeptides of the present invention. In another preferred embodiment, the present invention is directed to certain specific polynucleotides of SEQ ID NOS: 1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25, 27, 29, 31, 33, 35, 37, 39, 41, 43, 45, 47 and 49 that encode for the novel AAM polypeptides of SEQ ID NOS: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36, 38, 40, 42, 44, 46, 48 and 51, respectively. Preferred polynucleotides encode for a polypeptide of SEQ ID NO: 2, 6, 12, 16, 20, 24, 28, 30, 32, 34, 38, 44, 46 or 48; more preferably they encode a polypeptide of SEQ ID NO: 6, 12, 28, 34, 46 or 48; most preferably, they have a polypeptide of sequence of SEQ ID NO: 28 or 34.

[19] In a third aspect, the present invention is directed to a nucleic acid construct, a vector, or a host cell comprising a polynucleotide sequence encoding an AAM polypeptide of the present invention operatively linked to a promoter.

[20] In a fourth aspect, the present invention is directed to a method of making an AAM polypeptide of the present invention comprising (a) cultivating a host cell transformed with a nucleic acid sequence encoding an AAM polypeptide of the present invention under conditions suitable for production of the polypeptide; and (b) providing glucose to the cultivated host cells under conditions suitable for the production of  $\beta$ -alanine. The  $\beta$ -alanine may be optionally recovered from the cells.

[21] In a fifth aspect, the present invention is directed to a method of producing *b*-alanine comprising (a) cultivating a host cell transformed with a nucleic acid sequence encoding an AAM polypeptide of the present invention under conditions suitable for production of the polypeptide; and (b) providing glucose to the cultivated host cells under conditions suitable for the production of *b*-alanine. The *b*-alanine may be optionally recovered from the cells.

## BRIEF DESCRIPTION OF SEVERAL VIEWS OF THE DRAWINGS

[22] FIG. 1 shows the reversible reaction between alpha-alanine (*i.e.*, L-alanine or 2-aminopropionic acid) and **beta**-alanine (3-aminopropionic acid) that is catalyzed by alanine 2,3-aminomutase.

[23] FIG. 2 is a pathway for 3-hydroxypropionate (3-HP) synthesis from alpha-alanine, via beta-alanine as an intermediate.

[24] FIG. 3 is a 4036 bp expression vector (pCK110900-I Bla) of the present invention comprising a P15A origin of replication (P15A ori), a lacI repressor, a CAP binding site, a lac promoter (lac), a T7 ribosomal binding site (T7g10 RBS), and a chloramphenicol resistance gene (camR).

[25] FIGS. 4A-4J in combination provide an alignment chart of the amino acid sequences of four parental polypeptides that were used to produce the AAM of the present invention. The parental polypeptides were non-naturally occurring and derived in part from the KAM of *Clostrisium stricklandii* (SEQ ID NO: 53), *Porphyromonas gingivalis* (SEQ ID NO: 55), *Fusobacterium nucleatum* (SEQ ID NO: 57), and *Bacillus subtilis* (SEQ ID NO: 59), respectively. The sequences of two wild-type KAM are disclosed in SEQ ID NOS: 60 (P\_GI2529467\_G8\_AAB81159.1\_) and 61 (P\_GI2634361\_EMB\_CAB13860.1\_). A consensus sequence is also provided as SEQ ID NO: 62).

[26] The foregoing summary, as well as the following detailed description of certain embodiments of the present invention, will be better understood when read in conjunction with the appended drawings. For the purpose of illustrating the invention, there is shown in the drawings, certain embodiments. It should be understood, however, that the present invention is not limited to the arrangements and instrumentality shown in the attached drawings.

## DETAILED DESCRIPTION OF THE INVENTION

[27] The present invention has multiple aspects. In one aspect, the present invention is directed to a polypeptide having alanine 2,3-aminomutase (AAM) activity, preferably as measured by the assay of Example 8, and

- (a) having a polypeptide selected from the group consisting of SEQ ID NO: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36, 38, 40, 42, 44, 46, 48 and 51;
- (b) having an amino acid sequence which has at least 98% homology, with the amino acid sequence selected from the group consisting of SEQ ID NO: 2, 22, 28, 32, and 36;
- (c) having an amino acid sequence which has at least 99% homology, with the amino acid sequence selected from the group consisting of SEQ ID NO: 4, 6, 8, 12, 16, 24, 26, 30, 34 and 40;
- (d) being a polypeptide encoded by a nucleic acid sequence which hybridizes under high stringency conditions with either (i) the nucleotide sequence of SEQ ID NO: 1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25, 27, 29, 31, 33, 35, 37, 41, 43, 45, 47 or 49; (ii) a subsequence of (i) of at least 100 nucleotides, or (iii) a complementary strand of (i) or (ii) (J. Sambrook, E. F. Fritsch, and T. Maniatis, 1989, Molecular Cloning, A Laboratory Manual, 2d edition, Cold Spring Harbor, N.Y.); or
- (e) being a variant of the polypeptide of (d) comprising a substitution, deletion, and/or insertion of one to six amino acids therefrom and having AAM activity from about 1 to about 30  $\mu$ M  $\beta$ -alanine produced /hour 1 cell OD at pH 7.0-7.6, 25°C.

[28] Collectively, the polypeptides of (b) and (c) above are referred to herein as "homologous polypeptides." For purposes of the present invention, the degree of homology between two amino acid sequences is expressed as "percent homology," "percent identity," "% identity," "% identical," and "% identical" are used interchangeably herein to refer to the percent amino acid sequence identity that is obtained by ClustalW analysis (version W 1.8 available from European Bioinformatics Institute, Cambridge, UK), counting the number of identical matches in the alignment and dividing such number of identical matches by the length of the reference sequence, and using the following default ClustalW parameters to achieve slow/accurate pairwise optimal alignments – Gap Open Penalty:10; Gap Extension

Penalty:0.10; Protein weight matrix: Gonnet series; DNA weight matrix: IUB; Toggle Slow/Fast pairwise alignments = SLOW or FULL Alignment.

[29] AAM polypeptides are sensitive to oxygen and are preferably maintained and used in an oxygen deficient environment. If the AAM polypeptide becomes inactivated due to exposure to oxygen, it can be activated by anaerobic incubation with a sulphydryl compound for one hour at 37°C in accordance with the method described in Chirpich, et al., *Journal Biol. Chem.*, 245(7): 1778-1789 (1970), which is incorporated herein by reference in its entirety. AAM polypeptides of the present invention are preferably utilized in whole cell form (i.e., as a whole cell transformed with an AAM polynucleotide that is used under conditions such that the encoded AAM polypeptide is expressed in the cell) or alternatively, both isolated and utilized under anoxic conditions. AAM polypeptides of the present invention may be isolated, and optionally purified, under anaerobic conditions (e.g., under a nitrogen atmosphere) in accordance with the method described in Petrovich, et al., *Journal Biol. Chem.*, 266(12):7656-7660 (1991), which describes the isolation and purification of lysine-2,3-aminomutase and which is incorporated herein by reference in its entirety. As used herein, the term "anoxic" refers to oxygen deficient. The AAM polypeptides in whole cell form or as isolated enzymes may be lyophilized. In yet another embodiment, the present invention is directed to a composition comprising an AAM polypeptide as described herein (e.g., in whole cell form or as an isolated polypeptide) and a suitable carrier, typically a buffer, more typically an aqueous buffer solution having a pH from about 6.0 to about 8.0. It is also within the scope of the present invention that the aqueous buffered composition be lyophilized to provide a composition in a lyophilized form, wherein the composition is reconstituted by the addition of an aqueous based composition.

[30] In one embodiment, the present invention is also directed to an AAM polypeptide as described herein in isolated and purified form.

[31] In another embodiment, the present invention is directed to an AAM polypeptide as described herein in lyophilized form. Lyophilization is performed using standard lyophilization equipment. Typically, a solution containing the polypeptide is dispensed in an appropriate sized vial, frozen and placed under reduced

pressure to cause the water to evaporate, leaving the lyophilized (freeze-dried) polypeptide behind. Prior to use, the lyophilized polypeptide is reconstituted with distilled water or an appropriate buffer solution.

[32] In yet another embodiment, the present invention is directed to a composition comprising an AAM polypeptide as described herein and a suitable carrier, typically a buffer solution, more typically an aqueous buffer solution having a pH between 6.0 and 8.0. The composition may also be in a lyophilized form.

[33] The novel AAM polypeptides of the present invention have significantly enhanced AAM activity relative to the wild-type KAM polypeptides from which they are ultimately derived. By significantly enhanced AAM activity is meant that the AAM polypeptide of the present invention has an AAM activity within the range of about 1 to about 32  $\mu$ M  $\beta$ -alanine produced/hour 1 cell OD (units), preferably from about 10 to about 32 units, more preferably from about 20 to about 32 units; most preferably from about 25 to about 32 units.

[34] Table 1 provides a chart showing the AAM activities of the various AAM polypeptides of the present invention, identified by their clone number and SEQ ID NO. In Table 1, the OD<sub>600nm</sub> is reported at harvest after 5 hours (t=5) of incubation. Table 1 also reports the total  $\mu$ M of  $\beta$ -alanine produced after 5 hours per 1 cell OD. Finally, the last column of Table 1 reports the rate of  $\beta$ -alanine ( $\mu$ M) produced/hr /1 cell OD.

Table 1

Seq. ID No.	Harvest OD <sub>600nm</sub> t= 5	uM β-alanine produced at t=5/1 cell OD	Rate of β-alanine(uM) produced /hr 1 Cell OD
34	1.0	159.7	31.9
10	3.7	31.7	6.3
38	4.0	54.9	11.0
20	3.0	73.4	14.7
14	3.7	33.5	7.7
22	2.2	4.8	1.0
42	5.0	17.5	3.5
26	3.7	23.9	4.8
18	4.7	19.3	3.9
44	2.9	64.4	12.9
51	3.7	35.0	7.0
36	3.0	29.8	6.0
48	1.1	110.1	22.0
12	4.7	17.8	3.6
4	3.7	22.4	4.5
16	1.0	136.0	19.4
24	1.4	94.7	18.9
46	1.7	107.6	20.7
28	1.5	148.0	29.2
40	1.4	14.6	2.9
32	1.6	93.2	13.6
2	1.5	87.5	17.5
30	2.7	72.6	14.3
6	1.7	125.7	23.0

[35] Preferred AAM polypeptides of the present invention have an amino acid sequences of SEQ ID NOs: 2, 6, 12, 16, 20, 24, 28, 30, 32, 34, 38, 44, 46 or 48; more preferably they have an amino acid sequence of SEQ ID NOs: 6, 12, 28, 34, 46 or 48; most preferably, they have an amino acid sequence of SEQ ID NOs: 28 or 34.

[36] The ultimate grandparent molecule is the KAM of *Bacillus subtilis*, which had no detectable AAM activity. The DNA encoding this grandparent molecule was modified as described in WO 03/062173, entitled “Alanine 2,3-aminomutase,” to produce a polypeptide having a detectable alanine 2,3-aminomutase activity.

[37] In the present application, the applicants utilized as one parent molecule a polynucleotide of SEQ ID NO: 58, which encoded the 471 residue polypeptide of SEQ ID NO: 59 and which exhibited an AAM activity of approximately .001 U/mg (units/ mg of cell mass). The molecule of SEQ ID NO: 59 differs from the wild-type *B. subtilis* KAM (SEQ ID NO: 60), which had no detectable AAM activity, by having the following four (4) amino acid substitutions: L103M, M136V, Y140H and D339H.

[38] Other grandparent molecules utilized as starting materials in the present invention were the DNA sequences from other microorganisms (e.g., *Porphyromonas gingivalis*, *Fusobacterium nucleatum*, and *Clostridium sticklandii*) that encoded a KAM polypeptide. These DNA sequences were modified using standard techniques to introduce point substitutions that ultimately produced a KAM polypeptide that also had a detectable cross-reactivity with  $\alpha$ -alanine. One such parent molecule that was derived from *Porphyromonas gingivalis* is the polynucleotide of SEQ ID NO: 54 which encodes the 416 residue polypeptide of SEQ ID NO: 55. The parental polypeptide of SEQ ID NO: 55 differs from the wild-type *Porphyromonas gingivalis* KAM by having the following seven (7) amino acid substitutions: N19Y, E30K, L53P, H85Q, I192V, D331G, and M342T. Another such parent molecule that was derived from *F. nucleatum* is the polynucleotide of SEQ ID NO: 56 which encodes the 425 residue polypeptide of SEQ ID NO: 57.

[39] Yet another parent polynucleotide was derived by modification of the polynucleotide in *C. sticklandii* that encodes KAM. The resulting parental polynucleotide, which has a detectable cross-reactivity with  $\alpha$ -alanine, is the polynucleotide of SEQ ID NO: 52 which encodes the 416 residue polypeptide of SEQ ID NO: 53.

[40] The above described parental polypeptides of SEQ ID NOs: 53, 55, 57 and 58 are compared in the alignment chart of FIG. 4. From the alignment chart, it can be seen that the KAMs from *P. gingivalis*, *C. sticklandii*, and *F. nucleatum* are truncated at the N-terminus and at the C-terminus relative to the KAM from *B. subtilis*, while between the four species, about 40% of the residue positions in the central portion of the KAM polypeptide are conserved. Based upon the truncated species in the alignment chart of FIG. 4, it can be inferred that the first 8 amino acid residues at the

N-terminus of SEQ ID NO: 58 and the last 40 residues at the C-terminus of SEQ ID NO: 58 are not necessary for KAM activity, or the AAM activity that is derived therefrom. In FIG. 4, there is also provided a consensus sequence.

[41] The AAM polypeptide molecules of the present invention with their enhanced AAM activity were made by applying directed evolution techniques to the above-described parental molecules. These techniques are described in further detail herein.

[42] In yet another aspect, the present invention is directed to AAM polypeptides that have enhanced activity in coupled reactions.

[43] In another embodiment, the present invention is directed to an AAM a polypeptide encoded by a nucleic acid sequence which hybridizes under high stringency conditions with either (i) the nucleotide sequence of SEQ ID NO: 1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25, 27, 29, 31, 33, 35, 37, 41, 43, 45, 47 or 49; (ii) a subsequence of (i) of at least 100 nucleotides, or (iii) a complementary strand of (i) or (ii) (J. Sambrook, E. F. Fritsch, and T. Maniatis, 1989, Molecular Cloning, A Laboratory Manual, 2d edition, Cold Spring Harbor, N.Y.). For polynucleotides of at least 100 nucleotides in length, low to very high stringency conditions are defined as prehybridization and hybridization at 42°C in 5x SSPE, 0.3% SDS, 200 µg/ml sheared and denatured salmon sperm DNA, and either 25% formamide for low stringencies, 35% formamide for medium and medium-high stringencies, or 50% formamide for high and very high stringencies, following standard Southern blotting procedures.

[44] For polynucleotides of at least 100 nucleotides in length, the carrier material is finally washed three times each for 15 minutes using 2x SSC, 0.2% SDS at least at 50°C (low stringency), at least at 55°C (medium stringency), at least at 60°C. (medium-high stringency), at least at 65°C (high stringency), and at least at 70°C. (very high stringency).

[45] In another embodiment, the present invention is directed to a variant of the polypeptide of (d) comprising a substitution, deletion, and/or insertion of one to six amino acids therefrom and having AAM activity from about 1 to about 30 µM β-alanine produced /hour 1 cell OD at pH 7.0-7.6, 25°C, such as determined by the method of Example 8. Preferably, amino acid changes are of a minor nature, that is

conservative amino acid substitutions that do not significantly affect the folding and/or activity of the protein; small deletions, typically of one to six amino acids; small amino- or carboxyl-terminal extensions; a small linker peptide; or a small extension that facilitates purification by changing net charge or another function, such as a poly-histidine tract, an antigenic epitope or a binding domain.

[46] Examples of conservative substitutions are within the group of basic amino acids (arginine, lysine and histidine), acidic amino acids (glutamic acid and aspartic acid), polar amino acids (glutamine and asparagine), hydrophobic amino acids (leucine, isoleucine and valine), aromatic amino acids (phenylalanine, tryptophan and tyrosine), and small amino acids (glycine, alanine, serine, threonine, proline, cysteine and methionine). Amino acid substitutions, which do not generally alter the specific activity are known in the art and are described, for example, by H. Neurath and R. L. Hill, 1979, In, The Proteins, Academic Press, New York. The most commonly occurring exchanges are Ala/Ser, Val/Ile, Asp/Glu, Thr/Ser, Ala/Gly, Ala/Thr, Ser/Asn, Ala/Val, Ser/Gly, Tyr/Phe, Ala/Pro, Lys/Arg, Asp/Asn, Leu/Ile, Leu/Val, Ala/Glu, and Asp/Gly as well as these in reverse.

[47] In another embodiment, the present invention is directed to a fragment of (a), (b) or (c), as described above in the first paragraph of the Detailed Description, that has from about 1 to about 30  $\mu$ M  $\beta$ -alanine produced /hour 1 cell OD at pH 7.0-7.6, 25°C, such as determined by the method of Example 8. By the term "fragment" is meant that the polypeptide has a deletion of 1 to 8 amino acid residues from the N-terminus or 1-40 residues from the C-terminus, or both. Preferably, the deletion is 1 to 20 residues from the C-terminus, more preferably, the deletion is 1 to 10 residues from the C-terminus.

### Polynucleotides

[48] In its second aspect, the present invention is directed to a polynucleotide sequence that encodes for an AAM polypeptide of the present invention. Given the degeneracy of the genetic code, the present invention is also directed to any polynucleotide that encodes for the above referenced AAM polypeptides of the present invention. In its second aspect, the present invention is directed to a

polynucleotide sequence that encodes for the correspondingly referenced AAM polypeptide. Given the degeneracy of the genetic code, the present invention is also directed to any polynucleotide that encodes for the above referenced AAM polypeptides of the present invention. In a preferred embodiment, the present invention is directed to certain specific polynucleotides of SEQ ID NOS: 1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25, 27, 29, 31, 33, 35, 37, 39, 41, 43, 45, 47 and 49 that encode for the novel AAM polypeptides of SEQ ID NOS: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36, 38, 40, 42, 44, 46, 48 and 51, respectively. Preferred polynucleotides encode for a polypeptide of SEQ ID NO: 2, 6, 12, 16, 20, 24, 28, 30, 32, 34, 38, 44, 46 or 48; more preferably they encode a polypeptide of SEQ ID NO: 6, 12, 28, 34, 46 or 48; most preferably, they have a polypeptide of sequence of SEQ ID NO: 28 or 34.

[49] To make the improved AAM polypeptides of the present invention, one starts with one or more wild-type polynucleotides that encode a KAM polypeptide. The term "wild-type" polynucleotide means that the nucleic acid fragment does not comprise any mutations from the form isolated from nature. The term "wild-type" protein means that the protein will be active at a level of activity found in nature and typically will comprise the amino acid sequence as found in nature. Thus, the term "wild type" or "grand-parent sequence" indicates a starting or reference sequence prior to a manipulation of the invention.

[50] Suitable sources of wild-type KAM as a starting material to be improved is readily identified by screening genomic libraries for the KAM activity. A particularly suitable source of KAM is the *yodO* gene of *Bacillus sp.* bacteria as found in nature. Using the published KAM gene sequences for *B. subtilis* (e.g., WO 03 0623173 A2), primers for amplification of the genes from their respective gene libraries were created using conventional techniques. One such technique for isolating the KAM of *B. subtilis* is disclosed in Chen *et al.*, "A novel lysine 2,3-aminomutase encoded by the *yodO* gene of *Bacillus subtilis*: characterization on observation of organic radical intermediates," Biochem J. 348:539-549 (2000), which is incorporated herein by reference.

[51] The starting polynucleotides of SEQ ID NOs: 52, 54, 56 and 58 were obtained using the techniques disclosed in WO 03 0623173 A2 which is incorporated herein by reference for the disclosure of those techniques as recited in the examples therein. Specifically, WO 03 0623173 A2 discloses a *B. subtilis* wild-type lysine 2,3-aminomutase (KAM), and a mutated form thereof, which encodes an alanine 2,3-aminomutase (AAM). In addition, WO 03 0623173 A2 also discloses a *P. gingivalis* wild-type lysine 2,3-aminomutase (KAM) and a mutated form thereof, which encodes an alanine 2,3-aminomutase (AAM).

[52] Beginning with the polynucleotide of SEQ ID NO: 58, a non-naturally occurring and mutated and/or evolved enzyme, having unknown AAM activity is generated using any one of the well-known mutagenesis or directed evolution methods. See, e.g., Ling, et al., "Approaches to DNA mutagenesis: an overview," Anal. Biochem., 254(2):157-78 (1997); Dale, et al., "Oligonucleotide-directed random mutagenesis using the phosphorothioate method," Methods Mol. Biol., 57:369-74 (1996); Smith, "In vitro mutagenesis," Ann. Rev. Genet., 19:423-462 (1985); Botstein, et al., "Strategies and applications of in vitro mutagenesis," Science, 229:1193-1201 (1985); Carter, "Site-directed mutagenesis," Biochem. J., 237:1-7 (1986); Kramer, et al., "Point Mismatch Repair," Cell, 38:879-887 (1984); Wells, et al., "Cassette mutagenesis: an efficient method for generation of multiple mutations at defined sites," Gene, 34:315-323 (1985); Minshull, et al., "Protein evolution by molecular breeding," Current Opinion in Chemical Biology, 3:284-290 (1999); Christians, et al., "Directed evolution of thymidine kinase for AZT phosphorylation using DNA family shuffling," Nature Biotechnology, 17:259-264 (1999); Crameri, et al., "DNA shuffling of a family of genes from diverse species accelerates directed evolution," Nature, 391:288-291; Crameri, et al., "Molecular evolution of an arsenate detoxification pathway by DNA shuffling," Nature Biotechnology, 15:436-438 (1997); Zhang, et al., "Directed evolution of an effective fucosidase from a galactosidase by DNA shuffling and screening," Proceedings of the National Academy of Sciences, U.S.A., 94:454-4509; Crameri, et al., "Improved green fluorescent protein by molecular evolution using DNA shuffling," Nature Biotechnology, 14:315-319 (1996); Stemmer, "Rapid evolution of a protein in vitro by DNA shuffling," Nature, 370:389-391 (1994); Stemmer, "DNA shuffling by

random fragmentation and reassembly: *In vitro* recombination for molecular evolution," Proceedings of the National Academy of Sciences, U.S.A., 91:10747-10751 (1994); WO 95/22625; WO 97/0078; WO 97/35966; WO 98/27230; WO 00/42651; WO 01/75767 and U.S. Pat. 6,537,746 which issued to Arnold, *et al.* on March 25, 2003 and is entitled "Method for creating polynucleotide and polypeptide sequences."

[53] Any of these methods can be applied to generate AAM polynucleotides. To maximize any diversity, several of the above-described techniques can be used sequentially. Typically, a library of shuffled polynucleotides is created by one mutagenic or evolutionary technique and their expression products are screened to find the polypeptides having the highest AAM activity. Then, a second mutagenic or evolutionary technique is applied to polynucleotides encoding the most active polypeptides to create a second library, which in turn is screened for AAM activity by the same technique. The process of mutating and screening can be repeated as many times as needed, including the insertion of point mutations, to arrive at a polynucleotide that encodes a polypeptide with the desired activity, thermostability, or cofactor preference.

[54] Alternatively, polynucleotides and oligonucleotides of the invention can be prepared by standard solid-phase methods, according to known synthetic methods. Typically, fragments of up to about 100 bases are individually synthesized, then joined (e.g., by enzymatic or chemical ligation methods, or polymerase mediated methods) to form essentially any desired continuous sequence. For example, polynucleotides and oligonucleotides of the invention can be prepared by chemical synthesis using, e.g., the classical phosphoramidite method described by Beaucage *et al.* (1981) *Tetrahedron Letters* 22:1859-69, or the method described by Matthes *et al.* (1984) *EMBO J.* 3:801-05, e.g., as it is typically practiced in automated synthetic methods. According to the phosphoramidite method, oligonucleotides are synthesized, e.g., in an automatic DNA synthesizer, purified, annealed, ligated and cloned in appropriate vectors.

[55] In addition, essentially any nucleic acid can be custom ordered from any of a variety of commercial sources, such as The Midland Certified Reagent Company,

Midland, TX, The Great American Gene Company, Ramona, CA, ExpressGen Inc., Chicago, IL, Operon Technologies Inc., Alameda, CA, all of which have internet web sites, and many others. Similarly, peptides and antibodies can be custom ordered from any of a variety of sources, such as PeptidoGenic, HTI Bio-products, Inc., BMA Biomedicals Ltd. (U.K.), Bio.Synthesis, Inc., and many others.

[56] Polynucleotides may also be synthesized by well-known techniques as described in the technical literature. See, e.g., *Carruthers et al., Cold Spring Harbor Symp. Quant. Biol.* 47:411-418 (1982), and *Adams et al., J. Am. Chem. Soc.* 105:661 (1983). Double stranded DNA fragments may then be obtained either by synthesizing the complementary strand and annealing the strands together under appropriate conditions, or by adding the complementary strand using DNA polymerase with an appropriate primer sequence.

[57] General texts which describe molecular biological techniques useful herein, including mutagenesis, include Berger and Kimmel, Guide to Molecular Cloning Techniques, Methods in Enzymology, volume 152 Academic Press, Inc., San Diego, CA ("Berger"); Sambrook et al., Molecular Cloning - A Laboratory Manual (2nd Ed.), volumes 1-3, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, 1989 ("Sambrook"); and Current Protocols in Molecular Biology, F.M. Ausubel et al., eds., Current Protocols, a joint venture between Greene Publishing Associates, Inc. and John Wiley & Sons, Inc. (supplemented through 2000) ("Ausubel"). Examples of techniques sufficient to direct persons of skill through *in vitro* amplification methods, including the polymerase chain reaction (PCR) the ligase chain reaction (LCR), Q $\beta$ -replicase amplification and other RNA polymerase mediated techniques (e.g., NASBA) are found in Berger, Sambrook, and Ausubel, as well as Mullis et al., (1987) U.S. Patent No. 4,683,202; PCR Protocols A Guided to Methods and Applications (Innis et al., eds.) Academic Press Inc. San Diego, CA (1990); Arnheim & Levinson (October 1, 1990) Chemical and Engineering News 36-47; The Journal Of NIH Research (1991) 3:81-94; Kwoh et al. (1989) Proc. Natl. Acad. Sci. USA 86:1173; Guatelli et al. (1990) Proc. Natl. Acad. Sci. USA 87:1874; Lomell et al. (1989) J. Clin. Chem. 35:1826; Landegren et al., (1988) Science 241:1077-1080; Van Brunt (1990) Biotechnology 8:291-294; Wu and Wallace, (1989) Gene 4:560; Barringer et

*al.* (1990) *Gene* 89:117, and Sooknanan and Malek (1995) *Biotechnology* 13:563-564. Improved methods of cloning *in vitro* amplified nucleic acids are described in *Wallace et al.*, U.S. Pat. No. 5,426,039. Improved methods of amplifying large nucleic acids by PCR are summarized in *Cheng et al.* (1994) *Nature* 369:684-685 and the references therein, in which PCR amplicons of up to 40kb are generated. One of skill will appreciate that essentially any RNA can be converted into a double stranded DNA suitable for restriction digestion, PCR expansion and sequencing using reverse transcriptase and a polymerase. *See, Ausubel, Sambrook and Berger, all supra.*

[58] It will be appreciated by those skilled in the art due to the degeneracy of the genetic code, a multitude of nucleotide sequences encoding AAM polypeptides of the invention may be produced, some of which bear substantial identity to the nucleic acid sequences explicitly disclosed herein. It is also within the scope of the present invention that the polynucleotides encoding the AAM polypeptides of the present invention may be codon optimized for optimal production from the host organism selected for expression. Those having ordinary skill in the art will recognize that tables and other references providing codon preference information for a wide range of organisms are readily available. *See e.g., Henaut and Danchin, "Escherichia coli and Salmonella," Neidhardt, et al. Eds., ASM Press, Washington D.C., p. 2047-2066 (1996).*

[59] It is to be noted that expression in *E. coli* is different than in other organisms. For example, in the present invention, the codon (tgg) encodes Trp (W) for residue position 31 in the parent polypeptide of SEQ ID NO: 59. However, the corresponding codon for residue position 31 is "tga" in each of the progeny polynucleotides of SEQ ID NOs: 1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25, 27, 29, 31, 33, 35, 37, 39, 41, 43, 45, and 47 encoding for the AAM polypeptides of SEQ ID NOs: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36, 38, 40, 42, 44, 46, and 48, respectively. One skilled in the art recognizes that the codon "tga" is usually a stop (nonsense) codon. However, in the present expression system used in the  $\Delta$ panD *E. coli* strain, and under the selection conditions imposed, this codon is read through by the *E. coli* as a sense codon and is expressed, presumably as Trp (W). Others have reported that "tga" is the weakest stop codon for *E. coli* and that it is often read through as a sense codon

for Trp (W) in high expression. See e.g., Parker, J., "Errors and Alternatives in Reading the universal Genetic Code," *Microbiological Reviews*, 53(3): 273-298 (1989); Roth, J., "UGA Nonsense Mutations in *Salmonella typhimurium*," *J. of Bacteriology*, 102(2):467-475 (1970); and McBeath, G. and Kast, P., "UGA Read-Through Artifacts—When Popular Gene Expression Systems Need a Patch," *BioTechniques*, 24:789-794 (May 1998), which are incorporated herein by reference. Hence for expression in non-*E. coli* systems, it would be advantageous to alter the codon (tga) at residue position 31 to "tgg" which is the universal sense codon for Trp (W).

[60] In SEQ ID NO: 49, the codon encoding for residue 72 is "tag" which is read as a stop codon. However, two fragments are produced. The first fragment, having residues 1-71 of SEQ ID NO: 50, does not have any detectable AAM activity. The second fragment that is produced begins with residue 73 (Val) instead of the usual Met. This second fragment has 399 residues (SEQ ID NO: 51) and does have significant AAM activity (see Table 2) based upon the assay of Example 8. Thus, the first 72 residues at the N-terminus of the AAM polypeptide (based upon the consensus sequence or the parental KAM sequence from *B. subtilis*) are not absolutely necessary for AAM activity.

[61] In the present case, several round No. 1 libraries were created by applying a variety of mutagenic techniques to the polynucleotides of SEQ ID NOs: 52, 54, 56 and 58.

[62] In its third aspect, the present invention is directed to an expression vector and to a host cell comprising a polynucleotide of the present invention operatively linked to a control sequence. To obtain expression of the variant gene encoding an AAM polypeptide, the variant gene was first operatively linked to one or more heterologous regulatory sequences that control gene expression to create a nucleic acid construct, such as an expression vector or expression cassette. Thereafter, the resulting nucleic acid construct, such as an expression vector or expression cassette, was inserted into an appropriate host cell for ultimate expression of the AAM polypeptide encoded by the shuffled gene. A "nucleic acid construct" is defined herein as a nucleic acid molecule, either single- or double-stranded, which is isolated from a naturally

occurring gene or which has been modified to contain segments of nucleic acid combined and juxtaposed in a manner that would not otherwise exist in nature. Thus, in one aspect, the present invention is directed to a nucleic acid construct comprising a polynucleotide encoding an AAM polypeptide of the present invention.

[63] The term "nucleic acid construct" is synonymous with the term "expression cassette" when the nucleic acid construct contains all the control sequences required for expression of a coding sequence of the present invention. The term "coding sequence" is defined herein as a nucleic acid sequence, which directly specifies the amino acid sequence of its protein product. A coding sequence can include, but is not limited to, DNA, cDNA, and recombinant nucleic acid sequences.

[64] An isolated polynucleotide encoding an AAM polypeptide of the present invention may be manipulated in a variety of ways to provide for expression of the polypeptide. Manipulation of the isolated polynucleotide prior to its insertion into a vector may be desirable or necessary depending on the expression vector. The techniques for modifying polynucleotides and nucleic acid sequences utilizing recombinant DNA methods are well known in the art.

[65] The term "control sequence" is defined herein to include all components, which are necessary or advantageous for the expression of a polypeptide of the present invention. Each control sequence may be native or foreign to the nucleic acid sequence encoding the polypeptide. Such control sequences include, but are not limited to, a leader, polyadenylation sequence, propeptide sequence, promoter, signal peptide sequence, and transcription terminator. At a minimum, the control sequences include a promoter, and transcriptional and translational stop signals. The control sequences may be provided with linkers for the purpose of introducing specific restriction sites facilitating ligation of the control sequences with the coding region of the nucleic acid sequence encoding a polypeptide.

[66] The term "operably linked" is defined herein as a configuration in which a control sequence is appropriately placed at a position relative to the coding sequence of the DNA sequence such that the control sequence directs the expression of a polypeptide.

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[67] The control sequence may be an appropriate promoter sequence. The "promoter sequence" is a relatively short nucleic acid sequence that is recognized by a host cell for expression of the longer coding region that follows. The promoter sequence contains transcriptional control sequences, which mediate the expression of the polypeptide. The promoter may be any nucleic acid sequence which shows transcriptional activity in the host cell of choice including mutant, truncated, and hybrid promoters, and may be obtained from genes encoding extracellular or intracellular polypeptides either homologous or heterologous to the host cell.

[68] For bacterial host cells, suitable promoters for directing the transcription of the nucleic acid constructs of the present invention, include the promoters obtained from the *E. coli* lac operon, *Streptomyces coelicolor* agarase gene (dagA), *Bacillus subtilis* levansucrase gene (sacB), *Bacillus licheniformis* alpha-amylase gene (amyL), *Bacillus stearothermophilus* maltogenic amylase gene (amyM), *Bacillus amyloliquefaciens* alpha-amylase gene (amyQ), *Bacillus licheniformis* penicillinase gene (penP), *Bacillus subtilis* xylA and xylB genes, and prokaryotic beta-lactamase gene (Villa-Kamaroff et al., 1978, Proceedings of the National Academy of Sciences USA 75: 3727-3731), as well as the tac promoter (DeBoer et al., 1983, Proceedings of the National Academy of Sciences USA 80: 21-25). Further promoters are described in "Useful proteins from recombinant bacteria" in Scientific American, 1980, 242: 74-94; and in Sambrook et al., 1989, *supra*.

[69] For filamentous fungal host cells, suitable promoters for directing the transcription of the nucleic acid constructs of the present invention include promoters obtained from the genes for *Aspergillus oryzae* TAKA amylase, *Rhizomucor miehei* aspartic proteinase, *Aspergillus niger* neutral alpha-amylase, *Aspergillus niger* acid stable alpha-amylase, *Aspergillus niger* or *Aspergillus awamori* glucoamylase (glaA), *Rhizomucor miehei* lipase, *Aspergillus oryzae* alkaline protease, *Aspergillus oryzae* triose phosphate isomerase, *Aspergillus nidulans* acetamidase, and *Fusarium oxysporum* trypsin-like protease (WO 96/00787), as well as the NA2-tpi promoter (a hybrid of the promoters from the genes for *Aspergillus niger* neutral alpha-amylase and *Aspergillus oryzae* triose phosphate isomerase), and mutant, truncated, and hybrid promoters thereof.

[70] In a yeast host, useful promoters are obtained from the genes for *Saccharomyces cerevisiae* enolase (ENO-1), *Saccharomyces cerevisiae* galactokinase (GAL1), *Saccharomyces cerevisiae* alcohol dehydrogenase/glyceraldehyde-3-phosphate dehydrogenase (ADH2/GAP), and *Saccharomyces cerevisiae* 3-phosphoglycerate kinase. Other useful promoters for yeast host cells are described by Romanos et al., 1992, Yeast 8:423-488.

[71] The control sequence may also be a suitable transcription terminator sequence, a sequence recognized by a host cell to terminate transcription. The terminator sequence is operably linked to the 3' terminus of the nucleic acid sequence encoding the polypeptide. Any terminator, which is functional in the host cell of choice, may be used in the present invention.

[72] Preferred terminators for filamentous fungal host cells are obtained from the genes for *Aspergillus oryzae* TAKA amylase, *Aspergillus niger* glucoamylase, *Aspergillus nidulans* anthranilate synthase, *Aspergillus niger* alpha-glucosidase, and *Fusarium oxysporum* trypsin-like protease.

[73] Preferred terminators for yeast host cells are obtained from the genes for *Saccharomyces cerevisiae* enolase, *Saccharomyces cerevisiae* cytochrome C (CYC1), and *Saccharomyces cerevisiae* glyceraldehyde-3-phosphate dehydrogenase. Other useful terminators for yeast host cells are described by Romanos et al., 1992, *supra*.

[74] The control sequence may also be a suitable leader sequence, a nontranslated region of an mRNA which is important for translation by the host cell. The leader sequence is operably linked to the 5' terminus of the nucleic acid sequence encoding the polypeptide. Any leader sequence that is functional in the host cell of choice may be used in the present invention. Preferred leaders for filamentous fungal host cells are obtained from the genes for *Aspergillus oryzae* TAKA amylase and *Aspergillus nidulans* triose phosphate isomerase. Suitable leaders for yeast host cells are obtained from the genes for *Saccharomyces cerevisiae* enolase (ENO-1), *Saccharomyces cerevisiae* 3-phosphoglycerate kinase, *Saccharomyces cerevisiae* alpha-factor, and *Saccharomyces cerevisiae* alcohol dehydrogenase/glyceraldehyde-3-phosphate dehydrogenase (ADH2/GAP).

[75] The control sequence may also be a polyadenylation sequence, a sequence operably linked to the 3' terminus of the nucleic acid sequence and which, when transcribed, is recognized by the host cell as a signal to add polyadenosine residues to transcribed mRNA. Any polyadenylation sequence that is functional in the host cell of choice may be used in the present invention. Preferred polyadenylation sequences for filamentous fungal host cells are obtained from the genes for *Aspergillus oryzae* TAKA amylase, *Aspergillus niger* glucoamylase, *Aspergillus nidulans* anthranilate synthase, *Fusarium oxysporum* trypsin-like protease, and *Aspergillus niger* alpha-glucosidase. Useful polyadenylation sequences for yeast host cells are described by Guo and Sherman, 1995, Molecular Cellular Biology 15: 5983-5990.

[76] The control sequence may also be a signal peptide coding region that codes for an amino acid sequence linked to the amino terminus of a polypeptide and directs the encoded polypeptide into the cell's secretory pathway. The 5' end of the coding sequence of the nucleic acid sequence may inherently contain a signal peptide coding region naturally linked in translation reading frame with the segment of the coding region that encodes the secreted polypeptide. Alternatively, the 5' end of the coding sequence may contain a signal peptide coding region that is foreign to the coding sequence. The foreign signal peptide coding region may be required where the coding sequence does not naturally contain a signal peptide coding region.

[77] Alternatively, the foreign signal peptide coding region may simply replace the natural signal peptide coding region in order to enhance secretion of the polypeptide. However, any signal peptide coding region that directs the expressed polypeptide into the secretory pathway of a host cell of choice may be used in the present invention.

[78] Effective signal peptide coding regions for bacterial host cells are the signal peptide coding regions obtained from the genes for *Bacillus* NCIB 11837 maltogenic amylase, *Bacillus stearothermophilus* alpha-amylase, *Bacillus licheniformis* subtilisin, *Bacillus licheniformis* beta-lactamase, *Bacillus stearothermophilus* neutral proteases (nprT, nprS, nprM), and *Bacillus subtilis* prsA. Further signal peptides are described by Simonen and Palva, 1993, Microbiological Reviews 57: 109-137.

[79] Effective signal peptide coding regions for filamentous fungal host cells are the signal peptide coding regions obtained from the genes for *Aspergillus oryzae*

TAKA amylase, *Aspergillus niger* neutral amylase, *Aspergillus niger* glucoamylase, *Rhizomucor miehei* aspartic proteinase, *Humicola insolens* cellulase, and *Humicola lanuginosa* lipase.

[80] Useful signal peptides for yeast host cells are obtained from the genes for *Saccharomyces cerevisiae* alpha-factor and *Saccharomyces cerevisiae* invertase. Other useful signal peptide coding regions are described by Romanos *et al.*, 1992, *supra*.

[81] The control sequence may also be a propeptide coding region that codes for an amino acid sequence positioned at the amino terminus of a polypeptide. The resultant polypeptide is known as a proenzyme or propolypeptide (or a zymogen in some cases). A propolypeptide is generally inactive and can be converted to a mature active polypeptide by catalytic or autocatalytic cleavage of the propeptide from the propolypeptide. The propeptide coding region may be obtained from the genes for *Bacillus subtilis* alkaline protease (aprE), *Bacillus subtilis* neutral protease (nprT), *Saccharomyces cerevisiae* alpha-factor, *Rhizomucor miehei* aspartic proteinase, and *Myceliophthora thermophila* lactase (WO 95/33836).

[82] Where both signal peptide and propeptide regions are present at the amino terminus of a polypeptide, the propeptide region is positioned next to the amino terminus of a polypeptide and the signal peptide region is positioned next to the amino terminus of the propeptide region.

[83] It may also be desirable to add regulatory sequences, which allow the regulation of the expression of the polypeptide relative to the growth of the host cell. Examples of regulatory systems are those which cause the expression of the gene to be turned on or off in response to a chemical or physical stimulus, including the presence of a regulatory compound. In prokaryotic host cells, suitable regulatory sequences include the lac, tac, and trp operator systems. In yeast host cells, suitable regulatory systems include the ADH2 system or GAL1 system. In filamentous fungi, suitable regulatory sequences include the TAKA alpha-amylase promoter, *Aspergillus niger* glucoamylase promoter, and *Aspergillus oryzae* glucoamylase promoter.

[84] Other examples of regulatory sequences are those which allow for gene amplification. In eukaryotic systems, these include the dihydrofolate reductase gene, which is amplified in the presence of methotrexate, and the metallothionein genes, which are amplified with heavy metals. In these cases, the nucleic acid sequence encoding the AAM polypeptide of the present invention would be operably linked with the regulatory sequence.

### Expression Vectors

[85] In another aspect, the present invention is also directed to a recombinant expression vector comprising a polynucleotide of the present invention (which encodes an AAM polypeptide of the present invention), and one or more expression regulating regions. An expression regulating region includes a promoter, a terminator, a replication origin, etc., depending on the type of hosts into which they are to be introduced. The various nucleic acid and control sequences described above may be joined together to produce a recombinant expression vector which may include one or more convenient restriction sites to allow for insertion or substitution of the nucleic acid sequence encoding the polypeptide at such sites. Alternatively, the nucleic acid sequence of the present invention may be expressed by inserting the nucleic acid sequence or a nucleic acid construct comprising the sequence into an appropriate vector for expression. In creating the expression vector, the coding sequence is located in the vector so that the coding sequence is operably linked with the appropriate control sequences for expression.

[86] The recombinant expression vector may be any vector (e.g., a plasmid or virus), which can be conveniently subjected to recombinant DNA procedures and can bring about the expression of the polynucleotide sequence. The choice of the vector will typically depend on the compatibility of the vector with the host cell into which the vector is to be introduced. The vectors may be linear or closed circular plasmids.

[87] The expression vector may be an autonomously replicating vector, *i.e.*, a vector that, exists as an extrachromosomal entity, the replication of which is independent of chromosomal replication, *e.g.*, a plasmid, an extrachromosomal element, a minichromosome, or an artificial chromosome. The vector may contain

any means for assuring self-replication. Alternatively, the vector may be one which, when introduced into the host cell, is integrated into the genome and replicated together with the chromosome(s) into which it has been integrated. Furthermore, a single vector or plasmid or two or more vectors or plasmids which together contain the total DNA to be introduced into the genome of the host cell, or a transposon may be used.

[88] The expression vector of the present invention preferably contains one or more selectable markers, which permit easy selection of transformed cells. A selectable marker is a gene the product of which provides for biocide or viral resistance, resistance to heavy metals, prototrophy to auxotrophs, and the like. Examples of bacterial selectable markers are the *dal* genes from *Bacillus subtilis* or *Bacillus licheniformis*, or markers, which confer antibiotic resistance such as ampicillin, kanamycin, chloramphenicol (Example 1) or tetracycline resistance. Suitable markers for yeast host cells are ADE2, HIS3, LEU2, LYS2, MET3, TRP1, and URA3.

[89] Selectable markers for use in a filamentous fungal host cell include, but are not limited to, *amdS* (acetamidase), *argB* (ornithine carbamoyltransferase), *bar* (phosphinothricin acetyltransferase), *hph* (hygromycin phosphotransferase), *niaD* (nitrate reductase), *pyrG* (orotidine-5'-phosphate decarboxylase), (sulfate adenyltransferase), and *trpC* (anthranilate synthase), as well as equivalents thereof. Preferred for use in an *Aspergillus* cell are the *amdS* and *pyrG* genes of *Aspergillus nidulans* or *Aspergillus oryzae* and the *bar* gene of *Streptomyces hygroscopicus*.

[90] The vectors of the present invention preferably contain an element(s) that permits integration of the vector into the host cell's genome or autonomous replication of the vector in the cell independent of the genome. For integration into the host cell genome, the vector may rely on the nucleic acid sequence encoding the polypeptide or any other element of the vector for integration of the vector into the genome by homologous or nonhomologous recombination.

[91] Alternatively, the vector may contain additional nucleic acid sequences for directing integration by homologous recombination into the genome of the host cell. The additional nucleic acid sequences enable the vector to be integrated into the host cell genome at a precise location(s) in the chromosome(s). To increase the likelihood

of integration at a precise location, the integrational elements should preferably contain a sufficient number of nucleic acids, such as 100 to 10,000 base pairs, preferably 400 to 10,000 base pairs, and most preferably 800 to 10,000 base pairs, which are highly homologous with the corresponding target sequence to enhance the probability of homologous recombination. The integrational elements may be any sequence that is homologous with the target sequence in the genome of the host cell. Furthermore, the integrational elements may be non-encoding or encoding nucleic acid sequences. On the other hand, the vector may be integrated into the genome of the host cell by non-homologous recombination.

[92] For autonomous replication, the vector may further comprise an origin of replication enabling the vector to replicate autonomously in the host cell in question. Examples of bacterial origins of replication are P15A, pSC101, pMB1 and ColE1. Origins of replication of plasmids pBR322 (which has a pMB1 origin of replication), pUC19 (which has a ColE1 origin of replication), pACYC177 and pACYC184 (which have a P15A origin of replication), permit replication in *E. coli*; origins of replication for plasmids pUB110, pE194, pTA1060, or pAM beta.1 permit replication in *Bacillus*. Examples of origins of replication for use in a yeast host cell are the 2 micron origin of replication, ARS1, ARS4, the combination of ARS1 and CEN3, and the combination of ARS4 and CEN6. The origin of replication may be one having a mutation which makes its functioning temperature-sensitive in the host cell (see, e.g., Ehrlich, 1978, Proceedings of the National Academy of Sciences USA 75: 1433).

[93] More than one copy of a nucleic acid sequence of the present invention may be inserted into the host cell to increase production of the gene product. An increase in the copy number of the nucleic acid sequence can be obtained by integrating at least one additional copy of the sequence into the host cell genome or by including an amplifiable selectable marker gene with the nucleic acid sequence where cells containing amplified copies of the selectable marker gene, and thereby additional copies of the nucleic acid sequence, can be selected for by cultivating the cells in the presence of the appropriate selectable agent.

[94] The procedures used to ligate the elements described above to construct the recombinant nucleic acid construct and expression vectors of the present invention are

well known to one skilled in the art (see, e.g., J. Sambrook, E. F. Fritsch, and T. Maniatis, 1989, Molecular Cloning, A Laboratory Manual, 2d edition, Cold Spring Harbor, N.Y.).

[95] Many of the expression vectors for use in the present invention are commercially available. Suitable commercial expression vectors include p3xFLAGTM expression vectors from Sigma-Aldrich Chemicals, St. Louis MO., which includes a CMV promoter and hGH polyadenylation site for expression in mammalian host cells and a pBR322 origin of replication and ampicillin resistance markers for amplification in *E. coli*. Other suitable expression vectors are pBluescriptII SK(-) and pBK-CMV, which are commercially available from Stratagene, LaJolla CA, and plasmids that are derived from pBR322 (Gibco BRL), pUC (Gibco BRL), pREP4, pCEP4 (Invitrogen) or pPoly (Lathe et al., 1987, Gene 57, 193-201).

[96] Example 6 herein discloses the use of the expression vector pCK110900-I Bla, as shown in the vector map of FIG. 3.

### Host Cells

[97] Host cells for use in expressing the expression vectors of the present invention include but are not limited to, bacterial cells, such as *E. coli*, Streptomyces and *Salmonella typhimurium* cells; fungal cells, such as yeast cells (e.g., *Saccharomyces cerevisiae* or *Pichia pastoris* (ATCC Accession No. 201178)); insect cells such as *Drosophila* S2 and *Spodoptera* Sf9 cells; animal cells such as CHO, COS, 293, and Bowes melanoma cells; and plant cells. Appropriate culture mediums and conditions for the above-described host cells are well known in the art.

[98] By way of example, *Escherichia coli* W3110 was transformed by an expression vector for expressing the shuffled genes of the present invention. The expression vector was created by operatively linking a variant gene of the present invention to the *lac* promoter under control of the *lacI* repressor gene. The expression vector also contained the P15A origin of replication and the chloramphenicol resistance gene. The transformed *Escherichia coli* W3110 was cultured under appropriate culture medium containing chloramphenicol such that only transformed *E*

*coli* cells that expressed the expression vector survived. See e.g., Example 1. Purification

[99] Once the AAM polypeptides were expressed by the variant genes in *E. coli*, the polypeptides were purified from the cells and or the culture medium using any one or more of the well known techniques for protein purification, including lysozyme treatment, sonication, filtration, salting, ultra-centrifugation, affinity chromatography, and the like under strict anoxic conditions. Suitable solutions for high efficiency extraction of proteins from bacteria, such as *E. coli*, are commercially available under the trade name CelLytic B<sup>TM</sup> from Sigma-Aldrich of St. Louis MO. A suitable process for purifying AAM polypeptides sufficiently from cell lysate for applications in a chemical process is disclosed in the references: Chirpich, T. P. et al., J. Biol. Chem., 1970, 245, 1778-1789; and Petrovich, R. M. et al., J. Biol. Chem., 1991, 266, 7656-7660, both of which are incorporated herein by reference.

### Screening

[100] After several rounds of directed evolution were performed, the resulting libraries of exemplary AAM polypeptides were screened. Screening for transformed cells that express a polypeptide having AAM activity is, in general, a two-step process. First, one physically separates the cells and then determines which cells do and do not possess a desired property. Selection is a form of screening in which identification and physical separation are achieved simultaneously by expression of a selection marker, which, in some genetic circumstances, allows cells expressing the marker to survive while other cells die (or vice versa). Exemplary screening markers include luciferase,  $\beta$ -galactosidase, and green fluorescent protein. Selection markers include drug and toxin resistance genes, such as resistance to chloramphenicol, ampicillin and the like. Although spontaneous selection can and does occur in the course of natural evolution, in the present methods selection is performed by man.

[101] The AAM polynucleotides generated by the mutagenesis or directed evolution method are screened in accordance with the protocol described in Example 8 to identify those having enhanced activity that are suitable for inclusion as an improved AAM polypeptide of the present invention. In the process of Example 8, the

screening of clones from the expression libraries for enhanced AAM activity was performed by measuring the conversion of  $\alpha$ -alanine to  $\beta$ -alanine using liquid chromatography and mass spectrometry. Based upon the screening results, the AAM polypeptides of the present invention are listed in Table 2 below along with their residue changes and enhanced AAM activity relative to one parental AAM polypeptide, *i.e.*, the polypeptide of SEQ ID NO: 59.

**Table 2**

Seq. ID No.	Residue changes relative to parent SEQ ID NO: 59	Rate of $\beta$ -alanine(uM) produced /hr 1 Cell OD
34	I177L, I227M, G308R, I408L, F416S, D447G	31.9
10	I298V, G308R, F416S, D447G	6.3
38	D125N, I177L, T210S,	11.0
20	K2E, I307L,	14.7
14	K13E, L17R, L197P, I200T, M281V, F310S, F416S, D447G	7.7
22	Y72H, L118P, R145L, I220V, F240L, S250P, R311C, F416S, D447G	1.0
42	K19R, T99S, G308R, F416S, D447G	3.5
26	N80K, G308R, E319G, R325G, Q350R	4.8
18	Q32R, S74P, S113T, L118P, G308R, F416S, D447G	3.9
44	D79E, G308R, S329P, F393S, F414S, D445G, L453S,	12.9
51 (fragment)	A73V, G308R, Y331N, F416S, D447G	7.0
36	D79E, S93P, N132D, M281I, G308R, Y331N, F416S, D447G	6.0
48	K2E, M76I, D79E, T131A, L203P, G308R, Y331C, F416S, D447G	22.0
12	R38G, C134G, C141R, L203P, I280T, G308R, F416S, D447G	3.6
4	2KE, I220V, N237D, G308R, D360G, K361R, F416S, D447G	4.5

16	K13E, L17R, L197P, I200T, M281V, G308R, F310S, F416S, D447G	19.4
24	E23D, L43S, D124G, Y137H, K156E, G308R, D411G, F416S, D447G	18.9
46	W18R, M76I, D79E, V90A, M152T, I163T, S178P, V215G, G308R, V354A, F416S, D447G	20.7
28	E22G, Y71C, S74P, H108R, D187G, I244V, G308R, E396G, F416S, D447G, F454S	29.2
40	Y137H, G308R, D411G, F416S, D422V, D447G	2.9
32	H35R, D79E, K98T, T99S, N132S, S135P, E204G, K230R, G308R, F416S, D447G	13.6
2	W235R, S250P, C254R, D276G, G308R, Y380C, I381T, F416S, K440E, D447G	17.5
30	Q32R, N67S, H140R, G308R, F416S, D447G	14.3
6	E24G, M96I, E109G, G308R, F416S, D447G	23.0
8	G308R, S329P, F416S, D447G, L455S	14.7

[102] In Table 2 above, it is seen that the AAM polypeptides of the present invention have from 2 to 11 residue differences than their parent polypeptide of SEQ ID NO: 59, and very significant AAM activity as evidenced by the production of  $\beta$ -alanine in the assay of Example 8. In comparison,  $\beta$ -alanine was not detected for SEQ ID NO: 59 under the assay conditions used to test the AAM variants. However, some  $\beta$ -alanine production for parental SEQ ID NO: 59 was detected in a qualitative growth based complementation assay.

[103] Referring to Table 2 above, two preferred residue changes for the AAM polypeptides of the present invention relative to the parental sequence of SEQ ID NO: 59 are G308R and F416S. In those AAM polypeptides of the present invention that are at least 447 residues long, an additional preferred residue change is D447G relative to the parental sequence of SEQ ID NO: 59. Additional suitable residue

changes are G308K, F416M and D447L, A, I or V. Thus, in one aspect, the present invention is directed to an AAM polypeptide having at least 5 amino acid residue changes, typically 5-11 residue changes, relative to SEQ ID NO: 59 or a truncated fragment thereof as taught herein, the residue changes including from 1 to 3 residue changes selected from the group consisting of G308R, G308K, F416S, F416M, D447G, D447L, D447A, D447I and D447V.

[104] Based upon the AAM activity in Table 2, an especially preferred AAM polypeptide of the present invention is a polypeptide having 95% sequence homology with the polypeptide of SEQ ID NO: 34, more preferably 98% homology, most preferably 99% homology.

[105] The parental polypeptides of SEQ ID NOs: 53, 55 and 57 demonstrate that the residues 1-8 at the N-terminus and residues 434-473 at the C-terminus are not necessary for KAM or AAM activity. Likewise, the polypeptide fragment of SEQ ID NO: 51, which is a 399 residue expression product, discloses that the first 72 amino acids at the N-terminus relative to the parental clone of SEQ ID NO: 59 are not necessary for AAM activity. (See Table 2) Thus, it is also within the scope of the present invention that the polypeptides described herein include fragments thereof that lack from 1 to 72 residues from their N-terminus relative to the parental sequence of SEQ ID NO: 59, typically from 1 to 40 residues, more typically from 1-20 residues, most typically from 1 to 11 residues. It is also within the scope of the present invention that the above described N-terminal truncation be utilized in combination with a C-terminal truncation as described elsewhere herein.

[106] Only a very few ( $\leq 0.5\%$ ) of the mutations to the parental *B. subtilis* KAM (SEQ ID NO: 59) backbone were found to be beneficial. Specifically, for every 1000 clones screened, there occurred only 3-5 single point or double point mutations that were beneficial. In fact, some of the mutations were found to be detrimental.

[107] The first of the following two sets of sequences provides the sequence of the wild type *B. subtilis* lysine 2,3-aminomutase (KAM) polypeptides of the prior art, as deposited (GI\_2529467\_GB\_AAB81159.1). This sequence (SEQ ID NO: 60) was not used as a parent sequence but is provided only for purposes of comparison.

M K N K W Y K P K R H W K E I E L W K D V P E E K W N D W L W Q L T H T  
 V R T L D D L K K V I N L T E D E E E G V R I S T K T I P L N I T P Y Y A S L  
 M D P D N P R C P V R M Q S V P L S E E M H K T K Y D L E D P L H E D E D  
 S R V P G L T H R Y P D R V L F L V T N Q C S M Y C R Y C T R R R F S G Q I  
 G M G V P K K Q L D A A I A Y I R E T P E I R D C L I S G G D G L L I N D Q I  
 L E Y I L K E L R S I P H L E V I R I G T R A P V V F P Q R I T D H L C E I L K  
 K Y H P V W L N T H F N T S I E M T E E S V E A C E K L V N A G V P V G N  
 Q A V V L A G I N D S V P I M K K L M H D L V K I R V R P Y Y I Y Q C D L S  
 E G I G H F R A P V S K G L E I I E G L R G H T S G Y A V P T F V V D A P G G  
 G G K I A L Q P N Y V L S Q S P D K V I L R N F E G V I T S Y P E P E N Y I P  
 N Q A D A Y F E S V F P E T A D K K E P I G L S A I F A D K E V S F T P E N V  
 D R I K R R E A Y I A N P E H E T L K D R R E R R D Q L K E K K F L A Q Q K  
 K Q K E T E C G G D S S

[108] The second sequence in the set indicates the diversity of the AAM polypeptides of the present invention relative to the known wild-type *B. subtilis* KAM sequence by designating with the letter “X” followed by the residue number those residues in the Applicants’ AAM polypeptides that differ from those of wild-type *B. subtilis* KAM sequence:

M X<sub>2</sub> N K W Y K P K R H W X<sub>13</sub> E I E X<sub>17</sub> W X<sub>19</sub> D V P X<sub>23</sub> X<sub>24</sub> K W N D W L W  
 X<sub>32</sub> L T X<sub>35</sub> T V X<sub>38</sub> T L D D X<sub>43</sub> K K V I N L T E D E E E G V R I S T K T I P L  
 X<sub>67</sub> I T P X<sub>71</sub> X<sub>72</sub> X<sub>73</sub> X<sub>74</sub> L M D P X<sub>79</sub> X<sub>80</sub> P R C P V R M Q S V P L X<sub>93</sub> E E X<sub>96</sub> H  
 X<sub>98</sub> X<sub>99</sub> K Y D L E D P L X<sub>108</sub> X<sub>109</sub> D E D S X<sub>114</sub> V P G X<sub>118</sub> T H R Y P X<sub>124</sub> R V L F  
 L V T X<sub>132</sub> Q X<sub>134</sub> X<sub>135</sub> X<sub>136</sub> X<sub>137</sub> C R X<sub>140</sub> X<sub>141</sub> T R R X<sub>145</sub> F S G Q I G M G V P  
 X<sub>156</sub> K Q L D A A I A Y I R E T P E I R D C L I S G G D G L L I N X<sub>187</sub> Q I L E Y I  
 L K E X<sub>197</sub> R S X<sub>200</sub> P H X<sub>203</sub> X<sub>204</sub> V I R I G T R A P V V F P Q R I T D H X<sub>224</sub> C E I  
 L K X<sub>230</sub> X<sub>231</sub> H P V X<sub>235</sub> L X<sub>237</sub> T H X<sub>240</sub> N T S I E M T E E X<sub>250</sub> V E A X<sub>254</sub> E K L  
 V N A G V P V G N Q A V V L A G I N X<sub>276</sub> S V P X<sub>280</sub> X<sub>281</sub> K K L M H D L V K I  
 R V R P Y Y I Y Q C D L S E G X<sub>307</sub> X<sub>308</sub> H X<sub>310</sub> X<sub>311</sub> A P V S K G L X<sub>319</sub> I I E G L  
 R G H T X<sub>329</sub> G X<sub>331</sub> A V P T F V V X<sub>339</sub> A P G G G G K I A L X<sub>350</sub> P N Y V L S Q  
 S P X<sub>360</sub> K V I L R N F E G V I T S Y P E P E N X<sub>380</sub> X<sub>381</sub> P N Q A D A Y F E S V  
 X<sub>393</sub> P X<sub>395</sub> T A D K K E P I G L S A X<sub>408</sub> F A X<sub>411</sub> K E V S X<sub>416</sub> T P E N V X<sub>422</sub> R I  
 K R R E A Y I A N P E H E T L X<sub>440</sub> D R R E X<sub>445</sub> R X<sub>447</sub> Q L K E K K X<sub>454</sub> X<sub>455</sub> A  
 Q Q K K Q K E T E C G G D S S

The diversity of changes at various residue positions for the AAM polypeptides of the present invention are shown to the right of the arrow in Table 2 below and relative amino acid residues of wild-type KAM of *B. subtilis* (GI\_2529467\_GB\_AAB81159.1\_) (SEQ ID NO: 60) which are shown to the left of the arrow:

**Table 3**

X <sub>2</sub>	K→E
X <sub>13</sub> :	K→E
X <sub>17</sub> :	L→R
X <sub>19</sub> :	K→R
X <sub>23</sub> :	E→D, G
X <sub>24</sub> :	E→G
X <sub>32</sub> :	Q→R,
X <sub>35</sub> :	H→R
X <sub>38</sub> :	R→G
X <sub>43</sub> :	L→S
X <sub>67</sub> :	N→S
X <sub>71</sub> :	Y→C
X <sub>72</sub> :	Y→H, W
X <sub>73</sub> :	A→V
X <sub>74</sub> :	S→P
X <sub>79</sub> :	D→E
X <sub>80</sub> :	N→K
X <sub>93</sub> :	S→P
X <sub>96</sub> :	M→I
X <sub>98</sub> :	K→T
X <sub>99</sub> :	T→S
X <sub>108</sub> :	H→R
X <sub>109</sub> :	E→G
X <sub>114</sub> :	R→P
X <sub>118</sub> :	L→P
X <sub>124</sub> :	D→N
X <sub>132</sub> :	N→D, S
X <sub>134</sub> :	C→G
X <sub>135</sub> :	S→P
X <sub>136</sub> :	M→V
X <sub>137</sub> :	Y→H
X <sub>140</sub> :	Y→H
X <sub>141</sub> :	C→R
X <sub>145</sub> :	R→L
X <sub>156</sub> :	K→E
X <sub>187</sub> :	D→G
X <sub>197</sub> :	L→P
X <sub>200</sub> :	I→T
X <sub>203</sub> :	L→P
X <sub>204</sub> :	E→G
X <sub>224</sub> :	L→P
X <sub>230</sub> :	K→R
X <sub>231</sub> :	Y→H
X <sub>235</sub> :	W→R
X <sub>237</sub> :	N→D

X <sub>240</sub> :	F→L
X <sub>250</sub> :	S→P
X <sub>254</sub> :	C→Y, R
X <sub>276</sub> :	D→G
X <sub>280</sub> :	I→T
X <sub>281</sub> :	M→I, V
X <sub>307</sub> :	I→L
X <sub>308</sub> :	G→R
X <sub>310</sub> :	F→S
X <sub>311</sub> :	R→C
X <sub>319</sub> :	E→G
X <sub>329</sub> :	S→P
X <sub>331</sub> :	Y→N
X <sub>339</sub> :	D→H
X <sub>350</sub> :	Q→R
X <sub>360</sub> :	D→G
X <sub>361</sub> :	K→R
X <sub>380</sub> :	Y→C
X <sub>381</sub> :	I→T
X <sub>393</sub> :	F→S
X <sub>395</sub> :	E→G
X <sub>408</sub> :	I→L
X <sub>411</sub> :	D→G
X <sub>416</sub> :	F→S
X <sub>422</sub> :	D→V
X <sub>440</sub> :	K→E
X <sub>445</sub> :	R→K
X <sub>447</sub> :	D→G
X <sub>454</sub> :	F→S
X <sub>455</sub> :	L→S

[109] In a fourth aspect, the present invention is directed to a method of making an AAM a nucleic polypeptide of the present invention comprising (a) cultivating a host cell transformed with a nucleic acid sequence encoding an AAM polypeptide of the present invention under conditions suitable for production of the polypeptide; and (b) providing glucose to the cultivated host cells under conditions suitable for the production of β-alanine. The β-alanine may be optionally recovered from the cells.

**Example 1: Transformation protocol for *aam* libraries/ *ΔpanD* strain**

[110] A mutant *E. coli* strain - *ΔpanD*, derived from BW25113 which is described in Datsenko, K.A. and Wanner, B.L., Proc. Natl. Acad. Sci. USA 97:6640-6645 (2000)

was used as the host strain for screening of the *aam1* gene libraries. The protocol used to make the deletion is detailed in Example 4 of Cargill patent application WO 03/062173.

[111] Chemical competent *E. coli*  $\Delta$ *panD* was removed from -80°C frozen storage and thawed. Thereafter, it was kept on ice until used. An aliquot (100 $\mu$ l per transformation) was transferred into a sterile 1.5ml centrifuge tube. A KCM (5X) salt solution was added until the concentration in the aliquot was 1X. KCM consists of 700 mM KCl; 10 mM morpholinopropanesulphonic acid (MOPS) adjusted to pH 5.8. 1-5 $\mu$ l of the ligation mixture was added to the cells. The cells containing the ligation mixture were first incubated on ice for 30 minutes. The cells were heat shocked at 42°C for 1 min, and subsequently incubated on ice for 2 minutes. 500 $\mu$ l of SOC (Maniatis, T., Fritsch, E. F., and Sambrook, J. (1982) Molecular Cloning: A Laboratory Manual, 1st Ed., pp. A.2 and A.3, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY) was added to the cells, and the cells were incubated at 37°C for 1 hour with agitation. The cells were then centrifuged at 5000 rpm for 3 minutes, and the SOC was removed. The cell pellet was re-suspended in 500 $\mu$ l of M9 selection medium ((Maniatis, T., Fritsch, E. F., and Sambrook, J. (1982) Molecular Cloning: A Laboratory Manual, 1st Ed., pp. A.2 and A.3, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY) and incubated at 30°C for 2-4 hours with agitation. The cells were then plated onto M9 minimal agar medium supplemented with 1% mannose, 20 $\mu$ M iron citrate, 5.0 g/l  $\alpha$ -alanine, 0.1mM isopropyl- $\beta$ -D-thiogalactoside (IPTG) (Sigma Chemical Corp., St. Louis, MO), 50mM MOPS, 25mM bicarbonate, and 30 $\mu$ g/ml chloramphenicol. The plated cells were incubated at 30°C for 3 days or until colonies were of sufficient size to be picked using the Q-BOT™ robot colony picker (Genetix USA, Inc, Boston MA).

[112] In Round 2 of the transformation, the above procedure was followed except that the incubation temperature of the last two incubations in the procedure was increased to 37°C, and M9 minimal selection medium was not supplemented with  $\alpha$ -alanine (0 g/L  $\alpha$ -alanine).

**A. Alternate Transformation protocol for *aam* libraries/ *ΔpanD KIfldA* strain**

[113] A mutant *E. coli* strain *ΔpanD*, derived from BW25113 which is described in Datsenko, K.A. and Wanner, B.L., Proc. Natl. Acad. Sci. USA 97:6640-6645 (2000) is used as the host strain for screening of the *aam* gene libraries. The protocol used to make the deletion is detailed in Example 4 of International patent publication WO 03/062173. Optimally, a strain additionally having an increased expression of the flavodoxin (*fldA*) gene was used as the host strain for screening of the *aam* gene libraries, since increased flavodoxin enhances aminomutase activity when produced in *E. coli*. See USSN \_\_\_\_\_, by Cargill, Inc. (Liao, et al), filed October 14, 2005, entitled “Increasing the Activity of Radical S-Adenosyl Methionine (SAM) Enzymes” describes the production of β-alanine from cells that express AAM and overexpress flavodoxin at Examples 1-4, and these examples are incorporated herein by reference. This same application, USSN \_\_\_\_\_, by Cargill, Inc. (Liao, et al.) filed October 14, 2005, describes in Example 4 (incorporated herein) the construction of a strain of *E. coli* in which an artificial *P<sub>lac/ara</sub>* hybrid promoter was placed immediately upstream of the *fldA* gene. Strains carrying the artificial promoter before the *fldA* gene are designated *KIfldA*, where KI refers to “knock-in”).

[114] Competent cells of *E. coli* *ΔpanD KIfldA* are prepared either chemically or electrochemically using standard protocols. Competent *E. coli* *ΔpanD KIfldA* was removed from -80°C frozen storage and thawed. Thereafter, it was kept on ice until used. An aliquot (100μl per transformation) was transferred into a sterile 1.5ml centrifuge tube. A KCM (5X) salt solution was added until the concentration in the aliquot was 1X. KCM consists of 700 mM KCl; 10 mM morpholinopropanesulphonic acid (MOPS) adjusted to pH 5.8. 1-5μl of the ligation mixture was added to the cells. The cells containing the ligation mixture were first incubated on ice for 30 minutes. The cells were heat shocked at 42°C for 1 min, and subsequently incubated on ice for 2 minutes. 500μl of SOC (Maniatis, T., Fritsch, E. F., and Sambrook, J. (1982) Molecular Cloning: A Laboratory Manual, 1st Ed., pp. A.2 and A.3, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY) was added to the cells, and the cells were incubated at 37°C for 1 hour with agitation. The cells

were then centrifuged at 5000 rpm for 3 minutes, and the SOC was removed. Pellets were subsequently resuspended in a medium appropriate for either the complementation assay (Example 3) or the biotransformation assay (Example 4).

**Example 2: Cloning of *aam* genes into pCK110900 series vectors**

[115] The strategy employed for cloning the alanine aminomutase genes into an inducible expression system involved the isolation of the *aam* gene by PCR and cloning of the PCR fragment into the *SfiI* restriction sites downstream from a mutant *lac* promoter/operator system. Initially, PCR primers were designed to contain a nucleotide sequence that is specific to the 5' and 3' ends of the *aam* gene, as well as the Shine-Delgarno sequence of the ribosome-binding site, and the unique *SfiI* restriction sites. The gene was then amplified from a template, purified and digested with the restriction endonuclease *SfiI*. The restricted PCR fragment was purified using the QIAquick PCR purification kit (Qiagen), and cloned into the *SfiI* sites of the expression vector pCK110900-I Bla of FIG. 3 under the control of a *lac* promoter and *lacI* repressor gene. The expression vector also contained the P15a origin of replication and the chloramphenicol resistance gene. Shuffled *aam* gene libraries were cloned by the same method. Several clones were found that expressed an active alanine 2,3-aminomutase (as per the method of Example 8) and the synthetic genes were sequenced. A polynucleotide sequence designated BSAAM (SEQ ID NO: 58) - was used as a starting material for all further mutations and shuffling. BSAAM (SEQ ID NO: 58) has approximately 99.2% nucleotide identity with the wild-type *Bacillus subtilis* lysine aminomutase (GenBank Accession No. H10329).

**Example 3: Screening via the Tier 2a growth assay**

**Tier 2a growth Assay**

[116] The growth assay identifies variants capable of generating the essential metabolite AcetylCoA via  $\beta$ -alanine produced by AAM variants in the *E. coli*  $\Delta$ *panD* host strain. Growth is therefore a function of CoA production, and indirectly of AAM activity.

### A. Procedure

[117] AAM active clones from the tier 1 complementation assay were picked with a QBOT<sup>TM</sup> robot colony picker (Genetix USA, Inc., Boston MA) and inoculated into a 96-well master plate. The inoculums were grown in the 96 well master plate on a buffered minimal selection media (Na<sub>2</sub>HPO<sub>4</sub> 7H<sub>2</sub>O 12.8g/L; KH<sub>2</sub>PO<sub>4</sub> 3g/L; NaCl 0.5g/L; NH<sub>4</sub>Cl 1g/L; MgSO<sub>4</sub> 2mM; CaCl<sub>2</sub> 0.04mM; mannose 2%; IPTG 1mM; ferric citrate 20 uM; chloramphenicol 30 µg/ml; MOPS pH 7, 50mM; and sodium bicarbonate pH 9, 25mM) (hereinafter “MSM”) to which was added 0.1uM β-alanine and 0.5g/L α-alanine. Plates were covered using AirPore<sup>TM</sup> microporous tape (Qiagen, Inc.) and incubated at 25°C, 250 rpm, 85% humidity until cultures reached saturation, at which time glycerol was added to the cultures to a final concentration of 20-30%, and the plates stored at -80°C.

[118] Samples from a frozen master plate were arrayed into an “inoculum” plate containing buffered minimal selection media (MSM), as described above, further containing 0.5g/L α-alanine. The inoculum plates were covered with AirPore<sup>TM</sup> microporous tape (Qiagen, Inc.) and incubated at 25°C, 250 rpm, 85% humidity until cultures reached saturation.

[119] 15µl from the inoculum plate was inoculated into a 96-well “assay” plate containing 185µl of fresh MSM with 0.5g/L α-alanine. The assay plates were covered with AirPore<sup>TM</sup> microporous tape (Qiagen, Inc.) and a lid, and incubated at 25°C, 85% humidity, 250rpm. Measurements of OD at 600nm were made at discrete times for a period of approximately (~) 40hours.

### B. Data Analysis

[120] Since library variants exhibit unique growth profiles, it was preferable to calculate and compare growth rates (slopes) at three (3) different growth phases (early, mid and late) to identify all potentially improved variants. Clones that exhibit three (3) standard deviations above the plate average in any of the three (3) phases were designated as potentially improved variants and were retested in tier 2b for comparative ranking.

**Example 4: Screening via the Tier 2b growth assay**

[121] The stringency of the growth screen is increased in Tier 2b by excluding  $\alpha$ -alanine (the substrate for AAM) from the medium. Under these conditions, the cell relies on internal/cellular pools of  $\alpha$ -alanine to serve as a substrate for AAM, and subsequently, for cell growth. AAM variants capable of utilizing low, intracellular pools of  $\alpha$ -alanine might represent low  $K_M$  variants.

**A. Procedure**

[122] Samples from a frozen master plate were arrayed into an “inoculum” plate containing buffered minimal selection media (MSM), as described above, further containing 0.5g/L  $\alpha$ -alanine. The inoculum plates were covered using AirPore<sup>TM</sup> microporous tape and incubated at 25°C, 250 rpm, 85% humidity until cultures reached growth saturation.

[123] A TECAN<sup>TM</sup> Robotic Sample Processor (Columbus, Ohio) was used to remove 10 $\mu$ l of inoculum from each Tier 2a variant from the inoculum plates and seed it in replicates of 8 into each of the following:

96-well Assay plate containing 190 $\mu$ l of fresh MSM, 0.5g/L  $\alpha$ -alanine.

96-well Assay plate containing 190 $\mu$ l of fresh MSM, containing no  $\alpha$ -alanine.

The Assay plates were covered with AirPore<sup>TM</sup> microporous tape and a lid and grown at 25°C, 85% humidity, 250 rpm. Samples were collected at time points for approximately 3-4 days and the OD<sub>600nm</sub> was measured for each sample.

**B. Tier 2b Data Analysis**

[124] Variants were ranked by the following 3 criteria:

- i) Growth ratio equal to a final culture OD<sub>600</sub> on medium without  $\alpha$ -alanine/final culture OD<sub>600nm</sub> on medium containing  $\alpha$ -alanine;
- ii) Final culture OD<sub>600</sub>; and
- iii) Initial growth rates (in phase 1, from approximately 0-20 hour)

Clones with final culture OD<sub>600nm</sub> > 0.7 were retained.

Clones were then ranked based on the growth ratio of criteria (i). Any clones with a  $OD_{600nm} > 0.7$  were retained. However, clones that did not meet the above two criteria, but had a very good initial growth rate (iii) were also selected for further evaluation.

**Example 5: Screening via Tier 2c- PCR analysis**

The PCR screen identifies variants that contain the correct size gene in the expression vector prior to further screening for function. It excludes unstable gene variants that may have undergone deletions/truncations during the screening process.

**A. Procedure**

Potentially improved variants from frozen master plates were inoculated into a 96-microwell plate containing LB media with 1% glucose and 30  $\mu$ g/mL chloramphenicol. Cultures were grown at 25°C, 250 rpm, 85% humidity in plates covered with AirPore™ microporous tape (Qiagen, Inc.) until cultures reached saturation, approximately 2 days. 10  $\mu$ L of the culture was transferred to a PCR plate and boiled at 99°C for 10 minutes to disrupt the cells. Thereafter, 90  $\mu$ L of the following PCR Master Mix was added to the disrupted cells:

PCR Master Mix:

10 $\mu$ L	10X Taq Polymerase Buffer (QIAGEN, Valencia CA)
4 $\mu$ L	25 mM MgCl <sub>2</sub>
2 $\mu$ L	10 mM dNTPs
1.25 $\mu$ L	20 $\mu$ M primer – B <sub>forward</sub> (specific for BsAAM gene)
1.25 $\mu$ L	20 $\mu$ M primer – B <sub>reverse</sub> (specific for BsAAM gene)
1 $\mu$ L	5U/ $\mu$ L Taq polymerase (QIAGEN)
70.5 $\mu$ L	Sterile water
90 $\mu$ L	Total volume

The *Bacillus* specific primers used in the PCR reaction are as follows:

B-forward:

5'ccagcctggccataaggagatatacatatgaaaaacaaatggtataaac 3' SEQ ID NO: 63

B-reverse:

5' atggtgatggtgatggccagttggccttatgaagaatccctccgc 3' SEQ ID NO: 64

The amplification reaction was run for 30 cycles. The first cycle was run at 94°C for 1 minute. Thereafter, the extension procedure was performed for 29 cycles: 94.0°C for 1 minute; 55.0°C for 30 seconds; and 72.0°C for 1 minute. The final extension was performed at 72.0°C for 5 minutes. The products of the PCR reactions were analyzed by gel-electrophoresis on a 0.8% agarose gel.

**Example 6: Growth of AAM variants for β-alanine production (50 ml scale).**

**Cell selection method for identifying AAM activity.**

[125] To identify genes encoding polypeptides that can perform the alanine 2,3-aminomutase reaction, an efficient screen or selection for the desired activity is needed. Therefore, a selection method was developed by recognizing that *E. coli* uses beta-alanine for the synthesis of pantothenic acid, which in turn is a component of coenzyme A (CoA) and of acyl carrier protein (ACP). CoA and ACP are the predominant acyl group carriers in living organisms, and are essential for growth.

[126] In *E. coli*, the primary route to beta-alanine is from aspartate in a reaction catalyzed by aspartate decarboxylase (E.C. 4. 1. 1. 1), which is encoded by the *panD* gene. A functional deletion mutation of *panD* (shown as  $\Delta$ *panD*) results in beta-alanine auxotrophy and growth inhibition, which can be alleviated by the exogenous addition of pantothenate or beta-alanine, or by the production of beta-alanine from another source.

[127] Strain description: *E. coli*  $\Delta$ *panD* host (derived from BW25113, described in Datsenko, K.A. and Wanner, B.L., Proc. Natl. Acad. Sci. USA 97:6640-6645 (2000)), transformed with pCK110900-I Bla vector (low promoter strength resulting from mutated lac promoter sequence). The inoculum culture was grown in buffered minimal selection medium (MSM): M9 salts, pH 7.0-7.4, 50mM MOPs, pH 7.0, 25

mM sodium bicarbonate, pH 9.0, 1mM isopropyl- $\beta$ -D-thiogalactoside (IPTG), 30 $\mu$ g/ml chloramphenicol, 0.1g/L alanine, 5uM pyridoxine HCl, and 20uM ferric citrate. A 1:20 dilution of inoculum was used to inoculate 50ml of MSM medium described above. Cultures were incubated at 25°C, 250 rpm for approximately 3 days or until the culture reaches OD<sub>600nm</sub> ~1. Then,  $\alpha$ -alanine was added to the medium to a final concentration of 300 mM, and pantothenate was added to about 300uM. Incubation of the supplemented medium continued at 25°C, 250 rpm. Samples were removed from the medium for analysis at time points from t= 0 through t=5 hours following the addition of  $\alpha$ -alanine.

**Example 7: Method for extracting cells for  $\beta$ -alanine detection**

[128] Cells from the cultures of Example 6 were harvested by centrifugation of the cultures. The supernatant (spent media) was decanted and saved for further analysis (below). The cell pellets were washed with water. Pellets may be stored at -80°C for future extraction. The 50ml cell pellets (OD ~ 4.0) were re-suspended completely in a test tube in 0.9 ml water. The extraction volume for each sample was adjusted to this proportion according to the harvest OD<sub>600</sub>. An equal volume of methanol (-20°C) and 200  $\mu$ L of micro-glass beads was added and the mixture vortexed vigorously. Tubes containing the mixtures were placed on dry ice/EtOH, or in a -80°C freezer, for about 30 min. The frozen contents in the tube were thawed at room temperature and vortexed vigorously again, and centrifuged at maximum speed for about 10 minutes. The supernatants were filtered using 0.2–0.45 micron filter plates, or syringe filters.

[129] The spent medium was filtered using a 0.2-0.45 micron filter plate or syringe filter. The filtered spent medium was diluted 1:10 in -20°C methanol/water (final methanol concentration 50%).

[130] The  $\beta$ -alanine content of cell extract and spent media was analyzed by LC/MS/MS (Example 8).

For spent medium sample, the first minute was diverted to waste. The  $\beta$ -alanine peak arrived at approximately 2.0 minutes.

The assay can be scaled to 2ml, if only the spent media is analyzed.

**Example 8: Assay for  $\beta$ -alanine (LC/MS/MS)**

[131]  $\beta$ -alanine was determined using a combination of liquid chromatography and mass spectrometry. Suitable analytes were the cell extracts and spent media as prepared in Example 7.

[132] The liquid chromatography (LC) phase was performed using an ASTEC CHIROBIOTIC<sup>TM</sup> T 4.6 cm x 50 mm chiral LC column (Advanced Separation Technologies, Inc., Whippany, N.J. USA). The mobile phase consisted of two solutions: A: 0.25% aqueous acetic acid; and B: 0.25% (v/v) acetic acid in methanol. The elution was isocratic @ 0.6ml/minute.

[133] The mass spectrometer (MS) analysis was performed on a Micromass Ultima Triple Quad mass spectrometer, using the following tune parameters:

Capillary: 3.5 kV; cone: 20 V; hex 1: 15 V; aperture: 1.0V; source temp: 100°C; desolvation temp: 350°C; cone gas: 40 L/hr; desolvation gas: 500 L/h; low mass resolution(Q1): 12; high mass resolution (Q1): 12; ion energy (Q1): 0.1; collision cell entrance: -5; collision energy: 14; exit: 1; low mass resolution (Q2): 15 high mass resolution (Q2): 15; ion energy (Q2): 3.0; multiplier: 650 V.

**MS Method**

**Alanine transitions**

Analyte	Parent Ion (m/z)	Daughter Ion (m/z)	Dwell Time (sec)
$\alpha$ -alanine	90	44.7	0.1
$\beta$ -alanine	90	30.7	0.1
$\alpha$ -lysine	147	84.5	0.1
$\beta$ -lysine	147	70.5	0.1

The inter-channel delay was 0.1 seconds.

## CLAIMS

WHAT IS CLAIMED IS:

1. A polypeptide having alanine 2,3-aminomutase activity (hereinafter an "AAM polypeptide") and
  - (a) having an amino acid sequence selected from the group consisting of SEQ ID NO: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36, 38, 40, 42, 44, 46, 48 and 51;
  - (b) having an amino acid sequence which has at least 98% homology with the amino acid sequence selected from the group consisting of SEQ ID NO: 2, 22, 28, 32, and 36;
  - (c) having an amino acid sequence which has at least 99% homology with the amino acid sequence selected from the group consisting of SEQ ID NO: 4, 6, 8, 12, 16, 24, 26, 30, 34 and 40;
  - (d) being a polypeptide encoded by a nucleic acid sequence which hybridizes under high stringency conditions with either (i) the nucleotide sequence of SEQ ID NO: 1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25, 27, 29, 31, 33, 35, 37, 41, 43, 45, 47 or 49; (ii) a subsequence of (i) of at least 100 nucleotides, or (iii) a complementary strand of (i) or (ii); or
  - (e) being a variant of the polypeptide of (d) comprising a substitution, deletion, and/or insertion of one to six amino acids therefrom and having AAM activity from about 1 to about 30  $\mu$ M  $\beta$ -alanine produced /hour 1 cell OD at pH 7.0-7.6, 25°C.
2. The polypeptide of claim 1 having an amino acid sequence selected from the group consisting of SEQ ID NO: 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36, 38, 40, 42, 44, 46, 48 and 51.
3. The polypeptide of claim 1 having an amino acid sequence which has at least 98% homology with the amino acid sequence selected from the group consisting of SEQ ID NO: 2, 22, 28, 32, and 36.

4. The polypeptide of claim 1 having an amino acid sequence which has at least 99% homology with the amino acid sequence selected from the group consisting of SEQ ID NO: 4, 6, 8, 12, 16, 24, 26, 30, 34 and 40.

5. The polypeptide of claim 1 being a polypeptide encoded by a nucleic acid sequence which hybridizes under high stringency conditions with either (i) the nucleotide sequence of SEQ ID NO: 1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25, 27, 29, 31, 33, 35, 37, 41, 43, 45, 47 or 49; (ii) a subsequence of (i) of at least 100 nucleotides, or (iii) a complementary strand of (i) or (ii)

6. The polypeptide of claim 1 being a variant of the polypeptide of (d) comprising a substitution, deletion, and/or insertion of one to six amino acids therefrom and having AAM activity from about 1 to about 30  $\mu$ M  $\beta$ -alanine produced /hour 1/cell OD at pH 7.0-7.6, 25°C.

7. An AAM polypeptide having an amino acid sequence of SEQ ID NO: 2, 6, 12, 16, 20, 24, 28, 30, 32, 34, 38, 44, 46 or 48.

8. The AAM polypeptide of claim 7 having an amino acid sequence of SEQ ID NO: 6, 12, 28, 34, 46 or 48.

9. The AAM polypeptide of claim 8 having an amino acid sequence of SEQ ID NO: 28 or 34.

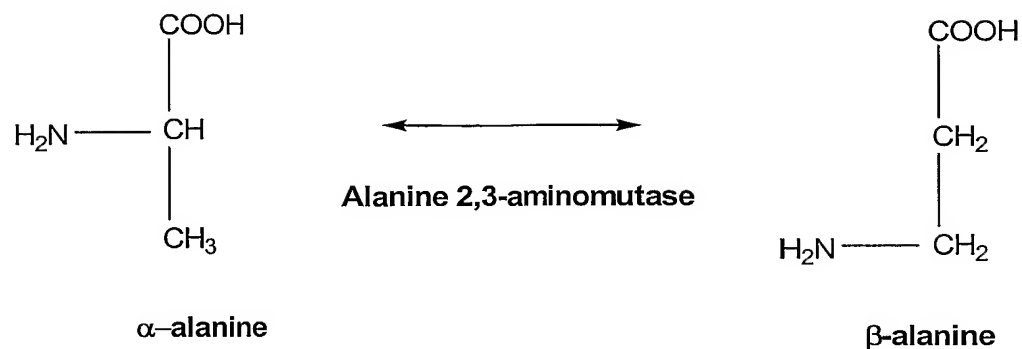
10. A polynucleotide encoding an AAM polypeptide of claim 1.

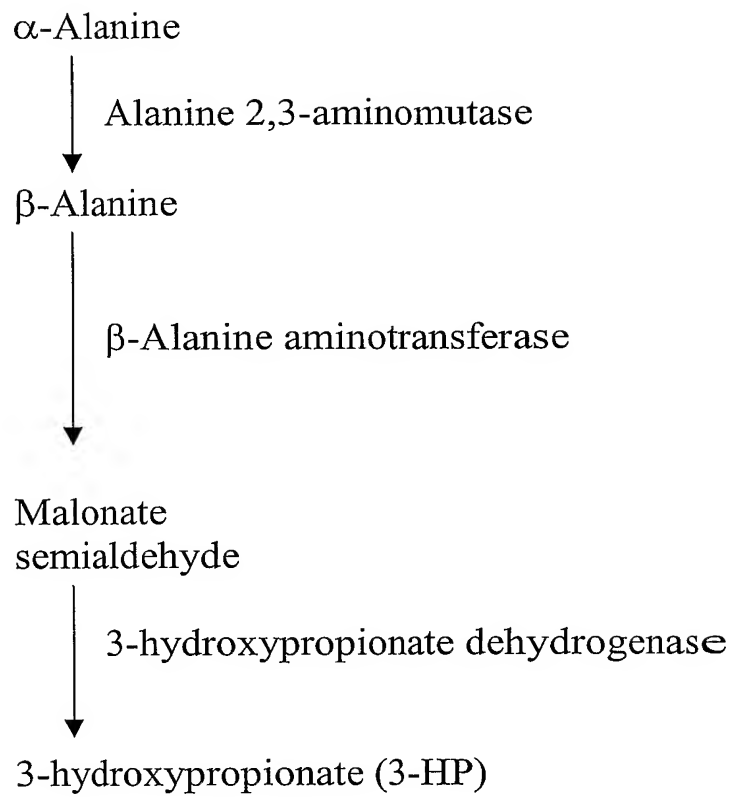
11. A polynucleotide encoding a polypeptide having AAM activity, said polynucleotide having SEQ ID NO: 1, 3, 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25, 27, 29, 31, 33, 35, 37, 41, 43, 45, 47 or 49.

12. An isolated and purified polynucleotide which encodes a polypeptide of claim 1.

13. An expression vector comprising a polynucleotide of claim 10 or 11 operatively linked to a promoter.

14. A host cell transformed to express a polynucleotide of claim 10.
15. A method of making an AAM polypeptide of claim 1, comprising (a) cultivating a host cell comprising a nucleic acid construct comprising a nucleic acid sequence encoding the AAM polypeptide under conditions suitable for production of the polypeptide; and (b) recovering the AAM polypeptide.
16. An AAM polypeptide of claim 1 in lyophilized form.
17. A composition comprising a polypeptide of claim 1 in a buffered medium.
18. An AAM polypeptide having from 5 to 11 amino acid residue changes relative to SEQ ID NO: 59 or a fragment thereof, the residue changes including from 1 to 3 residue changes selected from the group consisting of G308R, G308K, F416S, F416M, D447G, D447L, D447A, D447I and D447V.

**FIG. 1**

**FIG. 2**

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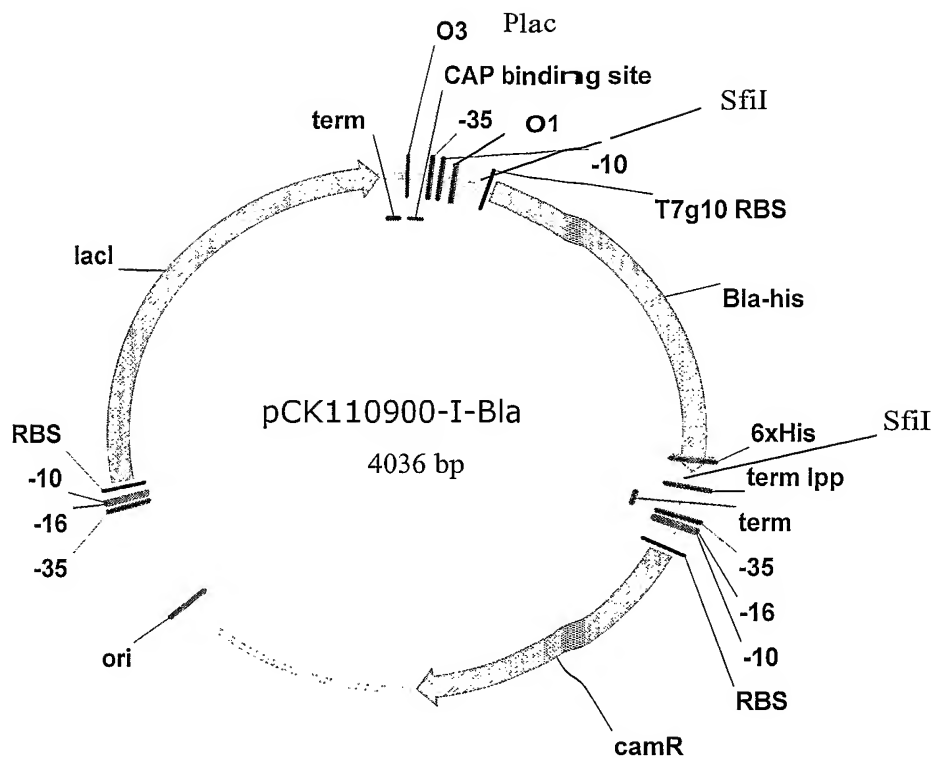


FIG. 3

4/8

SEQ ID NO:

1

50

P\_GI2529467\_G8\_AAB81159.1 60 (1)  
 MKNWYKPKRHWKEIELWKDVPEEKWNDWLWQLTHTVRLDDLKKVINLT  
 P\_GI2634361\_EMB\_CAB13860.1 61 (1)  
 MKNWYKPKRHWKEIELWKDVPEEKWNDWLWQLTHTVRLDDLKKVINLT  
 P\_S00701550 59 (1)  
 MKNWYKPKRHWKEIELWKDVPEEKWNDWLWQLTHTVRLDDLKKVINLT  
 P\_S00701551 53 (1) -----  
 MSLDKKFETHVSQEDWWDWKWQVRNRIKTVVEELKKYIPLT  
 P\_S00701552 55 (1) -----  
 MAESRRKYYEPDVTDEQWYDWHWQVLNRIKTLDDLKKYVTLIT  
 P\_S01032894 57 (1) -----  
 MNTVNTRKKFPPNVTDEEWNDWTWQVKNRNLKSVEDLEKYVDLS  
 Consensus 62 (1)  
 MKNWYKPKRHWKEIELWKDVPEEKWNDWLWQLTHTVRLDDLKKVINLT

**FIG. 4A**

51

100  
 P\_GI2529467\_G8\_AAB81159.1 (51)  
 EDEEEGVRISTKTIPLNITPYYASLMDPDNPRCPVRMOSVPLSEEMHKT  
 P\_GI2634361\_EMB\_CAB13860.1 (51)  
 EDEEEGVRISTKTIPLNITPYYASLMDPDNPRCPVRMOSVPLSEEMHKT  
 P\_S00701550 (51)  
 EDEEEGVRISTKTIPLNITPYYASLMDPDNPRCPVRMOSVPLSEEMHKT  
 P\_S00701551 (41)  
 PEEEGVKRCLDTLMAITPYYLSLIDVENPNPDPRKQAVPLSLELHRAA  
 P\_S00701552 (43)  
 AEEEGVKESPKVLRMAITPYYSLIDPENPNCPIRKQAIPTQQELVRAP  
 P\_S01032894 (44)  
 EEEEGVVRTLETLRMAITPFYFSLIDLNSDRCPIRKQAIPTIREIHQSD  
 Consensus (51)  
 EDEEEGVRISTKTIPLNITPYYASLMDPDNPRCPVRMOSVPLSEEMHKT

**FIG. 4B**

101  
 P\_GI2529467\_G8\_AAB81159.1 (101) YDLEDPIHEDEDSPVPGLTHRYPDRVTLFLVINVQCSMYCRYCTRRFSGQI  
 P\_GI2634361\_EMB\_CAB13860.1 (101) YDLEDPIHEDEDSPVPGLTHRYPDRVTLFLVINVQCSMYCRYCTRRFSGQI  
 P\_S00701550 (101) YDMDPDIHEDEDSPVPGLTHRYPDRVTLFLVINVQCSVYCRYCTRRFSGQI  
 P\_S00701551 (91) SDMDPDIHEDEDGDSPVPGLTHRYPDRVLLIMDQCSVYCRYCTRRFAGRT  
 P\_S00701552 (93) EDQVDESEDEDSPVPGLTHRYPDRVTLFLVINVQCSMYCRYCTRRFAGQK  
 P\_S01032894 (94) ADMIDPIHEDEDSPVPGLTHRYPDRVTLFLVINVQCSVYCRYCTRRFAGSS  
 Consensus (101) YDMDPDIHEDEDSPVPGLTHRYPDRVTLFLVINVQCSVYCRYCTRRFSGQI

FIG. 4C

150  
 P\_GI2529467\_G8\_AAB81159.1 (151) GMGVPIKKQLDAAIAYIRETPEIRDCLISGGDGLINDQILEYILKELRSI  
 P\_GI2634361\_EMB\_CAB13860.1 (151) GMGVPIKKQLDAAIAYIRETPEIRDCLISGGDGLINDQILEYILKELRSI  
 P\_S00701550 (151) GMGVPIKKQLDAAIAYIRETPEIRDCLISGGDGLINDQILEYILKELRSI  
 P\_S00701551 (141) DSAVDTKOIDAAEYIKNTEPQVRDVLLISGGDALLISDEKLEYTIRRLREI  
 P\_S00701552 (143) DASSPSPERIDRCIDYIANTPTVRDVLLISGGDALLVSDERLEYILKRLREV  
 P\_S01032894 (144) DGAMPMDRDKAYEYIAKTPQVRDVLLISGGDALLVSDERLEYILKRLRAI  
 Consensus (151) GMGVPIKKQLDAAIAYIRETPEIRDCLISGGDGLINDQILEYILKELRSI

FIG. 4D

201 P\_GI2529467\_G8\_AAB81159.1 (201)  
 P\_GI2634361\_EMB\_CAB13860.1 (201)  
 P\_S00701550 (201)  
 P\_S00701551 (191)  
 P\_S00701552 (193)  
 P\_S01032894 (194)  
 Consensus (201)

250 PHLEVIRIGTRAPVVFPQRITDHLCEILKKYHPVWLNTHTENTSIEMTTEES  
 PHLEVIRIGTRAPVVFPQRITDHLCEILKKYHPVWLNTHTENTSIEMTTEES  
 PHLEVIRIGTRAPVVFPQRITDHLCEILKKYHPVWLNTHTENTSIEMTTEES  
 PHLEVIRIGRSRVPPYMPORITPELVSMIRKYHPVWLNTIFNHPNEVTEEA  
 PHLEVIRIGSRTEVVLPORITPOLVDMKKYHPVWLNTIFNHPNEVTEEA  
 PHLEVIRIGSETPVLPORITPELCNMILKYHPVWLNTIFNHPQEVTPEA

FIG. 4E

300 251 VEACEKLVNAGVPVGNOAVLAGINDSVPIIMKKLMHDILVKIRVPRYYIYQ  
 (251) VEACEKLVNAGVPVGNOAVLAGINDSVPIIMKKLMHDILVKIRVPRYYIYQ  
 (251) VEACEKLVNAGVPVGNOAVLAGINDSVPIIMKKLMHDILVKIRVPRYYIYQ  
 (251) VEACEKLVNAGVPVGNOAVLAGINDSVPIIMKKLMHDILVKIRVPRYYIYQ  
 (241) KRACELLADAGIPLGNOQSVLIAQYNDCMHVMRKELVNDLVKIRVPRYYIYQ  
 (243) VEACERMANAGIPLGNOQTVLLRCINDCTHYAMRLYHILVKMRYMRYYIYV  
 (244) KKACEMLADAGVPLGNOTVHRLGINDSVPIIMKKLMHDILVKIRVPRYYIYQ  
 Consensus (251)

FIG. 4F

P_GI2529467_G8_AAB81159.1	(301)	CDLSEGIRHFRAPVSKGLELIEGLRHTSGYAVPTFVVDAPGGGKIALQ
P_GI2634361_EMB_CAB13860.1	(301)	CDLSEGIRHFRAPVSKGLELIEGLRHTSGYAVPTFVVDAPGGGKIALQ
P_S00701550	(301)	CDLSEGIRHFRAPVSKGLELIEGLRHTSGYAVPTFVVDAPGGGKIALQ
P_S00701551	(291)	CDLSEGIRHFRAPVSKGLELIEGLRHTSGYAVPTFVVDAPGGGKIALQ
P_S00701552	(293)	CDLSEGIRHFRAPVSKGLELIEGLRHTSGYAVPTFVVDAPGGGKIALQ
P_S01032894	(294)	CDLSEGIRHFRAPVSKGLELIEGLRHTSGYAVPTFVVDAPGGGKIALQ
Consensus	(301)	CDLSEGIRHFRAPVSKGLELIEGLRHTSGYAVPTFVVDAPGGGKIALQ

FIG. 4G

		400
351.		
P_GI2529467_G8_AAB81159.1	(351)	PNYVLQSPDKVILRNFEGVITSYPEPEVYIPNQADAYFESVFPETADKK
P_GI2634361_EMB_CAB13860.1	(351)	PNYVLQSPDKVILRNFEGVITSYPEPEVYIPNQADAYFESVFPETADKK
P_S00701550	(351)	PNYVLQSPDKVILRNFEGVITSYPEPEVYIPNQADAYFESVFPETADKK
P_S00701551	(341)	PNYVISONHNKYILRNFEGVITTYDRBDHYTFHCDVCTGKT-----NV
P_S00701552	(343)	PNYVVSQSPRHVYLRNFEGVITTYEPENYHEECDCEDCRAG-----K
P_S01032894	(344)	POYVISOSPHRAVILRNFEGVITTYEPENYTHEPCYDEEKKFEK-----MY
Consensus	(351)	PNYVLQSPSPDKVILRNFEGVITSYPEPEVYIPNQADAYFESVFPETADKK

FIG. 4H

450

P_GI2529467_G8_AAB81159.1	(401)	EPIGLSAILADKEVSETENVDRIKRREAYIANPEHEETLKDERRDOLK
P_GI2634361_EMB_CAB13860.1	(401)	EPIGLSAILADKEVSETENVDRIKRREAYIANPEHEETLKDERRDOLK
P_S00701550	(401)	EPIGLSAILADKEVSETENVDRIKRREAYIANPEHEETLKDERRDOLK
P_S00701551	(386)	HKIVGVAGLNLGETATIEDEGLERKORGHH-----
P_S00701552	(386)	HKEGVAAATSGQOLATEPSDEARKKRFIDKN-----
P_S01032894	(389)	EISGVYMLDEGLEMSLEPSHILARHERNIKKRAEAGKK-----
Consensus	(401)	EPIGLSAILADKEVSETENVDRIKRREAYIANPEHEETLKDERRDOLK

**FIG. 4I**

P_GI2529467_G8_AAB81159.1	(451)	EKKFLAQQQKKQETECGGDSS-----
P_S00701550	(451)	EKKFLAQQQKKQETECGGDSS-----
P_S00701551	(415)	-----
P_S00701552	(417)	-----
P_S01032894	(426)	-----
Consensus	(451)	EKKFLAQQQKKQETECGGDSS-----

**FIG. 4J**

-1-

SEQUENCE LISTING

<110> Chatterjee, Ranjini  
Chien, Michelle  
Louie, Susan  
Mitchell, Ken  
Fox, Richard

<120> Improved Alanine 2,3-Aminomutases and Related Polynucleotides

<130> 0359.210WO/15686WO02

<160> 64

<170> PatentIn version 3.3

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Leu	Trp	Lys	Asp	Val	Pro	Glu	Glu	Lys	Trp	Asn	Asp	Trp	Leu	Trp	Gln
				20				25					30		

Leu	Thr	His	Thr	Val	Arg	Thr	Leu	Asp	Asp	Leu	Lys	Lys	Val	Ile	Asn
				35				40				45			

Leu	Thr	Glu	Asp	Glu	Glu	Gly	Val	Arg	Ile	Ser	Thr	Lys	Thr	Ile	
				50			55		60						

Pro	Leu	Asn	Ile	Thr	Pro	Tyr	Tyr	Ala	Ser	Leu	Met	Asp	Pro	Asp	Asn
65				70				75			80				

Pro	Arg	Cys	Pro	Val	Arg	Met	Gln	Ser	Val	Pro	Leu	Ser	Glu	Glu	Met
				85				90				95			

His	Lys	Thr	Lys	Tyr	Asp	Met	Glu	Asp	Pro	Leu	His	Glu	Asp	Glu	Asp
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Ser	Pro	Val	Pro	Gly	Leu	Thr	His	Arg	Tyr	Pro	Asp	Arg	Val	Leu	Phe
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Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Arg Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Pro Val Glu Ala Arg Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Gly Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

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Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Cys Thr Pro Asn Gln  
 370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
 385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
 405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
 420 425 430

Asn Pro Glu His Glu Thr Leu Glu Asp Arg Arg Glu Lys Arg Gly Gln  
 435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
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Glu Cys Gly Gly Asp Ser Ser  
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 gatgggctgc tcatcaacga ccaaatttttta gaatataattt taaaagagct ggcgcagcatt 360  
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 480  
 540  
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<220>  
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Leu Trp Lys Asp Val Pro Glu Glu Lys	Trp Asn Asp Trp Leu Trp Gln	
20	25	30

Leu Thr His Thr Val Arg Thr Leu Asp	Asp Leu Lys Lys Val Ile Asn	
35	40	45

Leu Thr Glu Asp Glu Glu Glu Gly Val	Arg Ile Ser Thr Lys Thr Ile	
50	55	60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala	Ser Leu Met Asp Pro Asp Asn		
65	70	75	80

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Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Val Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asp Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

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Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
 305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
 325 330 335

Val Val His Ala Pro Gly Gly Gly Lys Ile Ala Leu Gln Pro Asn  
 340 345 350

Tyr Val Leu Ser Gln Ser Pro Gly Arg Val Ile Leu Arg Asn Phe Glu  
 355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
 370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
 385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
 405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
 420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
 435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
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Glu Cys Gly Gly Asp Ser Ser  
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 gatgatttaa agaaagtcat taatctgacc gaggatgaag aggaaggcgt ccgtattct 180

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cgctatcccg	accgtgtgct	gtttcttgc	acgaatcaat	gttctgtgt	ctgcccac	42	0	
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Leu Trp Lys Asp Val Pro Glu Gly Lys Trp Asn Asp Trp Leu Trp Gln  
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Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Ile  
85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Gly Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

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Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
450 455 460

Glu Cys Gly Gly Asp Ser Ser  
465 470

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gatgatttaa agaaaagt~~c~~at taatctgacc gaggatgaag aggaaggcgt ccgtatttct 180  
accaaaaacga tcccctt~~a~~aa tattacacca tactatgcga gcttaatgga tccagaaaaac 240  
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<210> 8  
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<213> Artificial Sequence

<220>  
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Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln  
20 25 30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Glu Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

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Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Pro  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Pro Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Ser Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
 435 440 445

Leu Lys Glu Lys Lys Phe Ser Ala Gln Gln Lys Lys Gln Lys Glu Thr  
 450 455 460

Glu Cys Gly Gly Asp Ser Ser  
 465 470

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 gatgatttaa agaaaagtcat taatctgacc gaggatgaag aggaaggcgt ccgtatttct 180  
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 tgcacacgcc ggcgttttc cggacaaaatc ggaatggcg tccccaaaaa acagcttgat 480  
 gctgcaatttgc ttatatccg ggaaacaccc gaaatcccg attgttaat ttcaggcggt 540  
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 accgatcatc tgtgcgagat attgaaaaaa tatcatccg tctggctgaa cacccatttt 720  
 aacacaagca tcgaaatgac agaagaatcc gttgaggcat gtgaaaagct ggtgaacgcg 780  
 ggagtgcgg tcggaaatca ggctgtcgta ttagcaggta ttaatgattc ggttccaatt 840  
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 attgaagggc tgagaggtca tacctcaggc tatgcgggtc ctactttgt cgttcacgca 1020  
 ccaggcggag gaggtaaaat cgcccctgcag ccgaactatg tcctgtctca aagtccctgac 1080

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aaagtatct taagaaattt tgaaggtgtg attacgtcat atccggaacc agagaattat 1140  
 atccccaaatc aggcagacgc ctattttag tccgtttcc ctgaaaccgc tgacaaaaag 1200  
 gagccgatcg ggctgagtgc cattttgct gacaaagaag tttcgtctac acctgaaaat 1260  
 gtagacagaa tcaaacggcg tgaggcatac atcgcaaatc cggagcatga aacattaaaa 1320  
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<210> 10  
 <211> 471  
 <212> PRT  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct  
 <400> 10

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu  
 1 5 10 15

Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln  
 20 25 30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
 35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
 50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Asn  
 65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
 85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
 100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
 115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
 130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Val Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

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Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
 370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
 385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
 405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
 420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
 435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
 450 455 460

Glu Cys Gly Gly Asp Ser Ser  
 465 470

<210> 11  
 <211> 1416  
 <212> DNA  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<400> 11  
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 gtcccggaag agaaatggaa cgattggctt tgacagctga cacacactgt aggaacgtta  
 gatgatttaa agaaagtcat caatctgacc gaggatgaag aggaaggcgt ccgtatttct 120  
 accaaaaacga tccccttaaa tattacacct tactatgctt cttaatgga ccccgacaat  
 ccgagatgcc cggtagcgtat gcagtctgtc ccgtttctg aagaaatgca caaaacaaaa 180  
 tacgatatgg aagacccgct tcatgaggat gaagattcac cggtaccgg tctgacacac  
 cgctatcccc accgtgtgct gtttcttgc acgaatcaag gttccgtgta ctgcccccac 240  
 cgcacacgcc ggcgcctttc cggacaaatc ggaatggcgc tccccaaaaa acagcttgat  
 gctgcaattt cttatatccg ggaaacaccc gaaatccgcg atttttaat ttcaggcggt 300  
 gatgggctgc tcatcaacga ccaaatttttta gaatatattt taaaagagct ggcgcagcatt  
 ccgcattccgg aagtcatccg catcggaaaca cgtgctcccg tcgtttccc gcagcgcatt 360  
 420  
 480  
 540  
 600  
 660

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accgatcatc	tgtgcgagat	attgaaaaaa	tatcatccgg	tctggctgaa	cacccatttt	720
aacacaagca	tcgaaatgac	agaagaatcc	gttgaggcat	gtgaaaagct	ggtgaacgcg	780
ggagtgccgg	tcggaaatca	ggctgtcgta	ttagcaggtta	ttaatgattc	ggttccaact	840
atgaaaaaagc	tcatgcatga	cttggtaaaa	atcagagtcc	gtccttatta	tatcaccaa	900
tgtgatctgt	cagaaggaat	aaggcatttc	cgtgctcctg	tttccaaagg	tttggagatc	960
attgaagggc	tgagaggcca	tacctcaggc	tatgcggttc	ctaccttgc	cgttcacgca	1020
ccaggcggag	gaggtaaaat	cgcgcgcag	ccgaactatg	tcctgtctca	aagtccctgac	1080
aaagtgatct	taagaaattt	tgaaggtgtg	attacgtcat	atccggaacc	agagaattat	1140
atccccaaatc	aggcagacgc	ctatttgag	tccgtttcc	ctgaaaccgc	tgacaaaaag	1200
gagccgatcg	ggctgagtgc	cattttgct	gacaaagaag	tttcgtctac	acctgaaaat	1260
gtagacagaa	tcaaacggcg	tgaggcatac	atcgcaaatc	cggagcatga	aacattaaaa	1320
gatcggcgtg	agaaaagagg	tcagctaaa	gaaaagaaaat	tttggcgca	gcagaaaaaa	1380
cagaaagaga	ctgaatgcgg	aggggattct	tcataa			1416

<210> 12

<211> 471

<212> PRT

<213> Artificial Sequence

<220>

<223> Synthetic Construct

<400> 12

Met	Lys	Asn	Lys	Trp	Tyr	Lys	Pro	Lys	Arg	His	Trp	Lys	Glu	Ile	Glu
1				5				10				15			

Leu	Trp	Lys	Asp	Val	Pro	Glu	Glu	Lys	Trp	Asn	Asp	Trp	Leu	Trp	Gln
				20				25				30			

Leu	Thr	His	Thr	Val	Gly	Thr	Leu	Asp	Asp	Leu	Lys	Lys	Val	Ile	Asn
				35				40				45			

Leu	Thr	Glu	Asp	Glu	Glu	Glu	Gly	Val	Arg	Ile	Ser	Thr	Lys	Thr	Ile
				50				55				60			

Pro	Leu	Asn	Ile	Thr	Pro	Tyr	Tyr	Ala	Ser	Leu	Met	Asp	Pro	Asp	Asn
65				70					75			80			

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Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Gly Ser Val Tyr Cys Arg His Arg Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Pro Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Thr Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

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Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
 325 330 335

Val Val His Ala Pro Gly Gly Lys Ile Ala Leu Gln Pro Asn  
 340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
 355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
 370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
 385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
 405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
 420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
 435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
 450 455 460

Glu Cys Gly Gly Asp Ser Ser  
 465 470

<210> 13

<211> 1416

<212> DNA

<213> Artificial Sequence

<220>

<223> Synthetic Construct

<400> 13

atgaaaaaca aatggataaa accgaaacgg cattgggagg agatcgagcg atggaaggac 60

gttccggaaag agaaatggaa cgattggctt tgacagctga cacacactgt aagaacgtta 120

gatgatttaa agaaagtcat taatctgacc gagggatgaag aggaaggcgt ccgtatttct 180

acccaaaacga tcccttaaaa tattacacct tactatgctt ccttaatgga ccccgacaat 240

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<210> 14  
<211> 471  
<212> PRT  
<213> Artificial Sequence

<220>  
<223> Synthetic Construct

<400> 14

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Glu Glu Ile Glu  
 1 5 10 15

Arg Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln  
20 25 30

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Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Pro Arg Ser Thr Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

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Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Val Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Ser Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
450 455 460

Glu Cys Gly Gly Asp Ser Ser  
465 470

<210> 15  
<211> 1416

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&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; Synthetic Construct

&lt;400&gt; 15

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gttccggaag agaaatggaa cgattggctt tgacagctga cacacactgt aagaacgtta	12 0
gatgatttaa agaaagtcat taatctgacc gaggatgaag aggaaggcgt ccgtatttct	18 0
accaaaaacga tccccttaaa tattacacct tactatgctt ccttaatgga ccccgacaaat	24 0
ccgagatgcc cggtagcat gcagtctgtg ccgccttctg aagaaatgca caaaacaaaa	30 0
tacgatatgg aagacccgct tcatgaggat gaagattcac cggtagccgg tctgacacac	36 0
cgctatcccg accgtgtgct gtttcttgc acgaatcaat gttccgtgta ctgccgccac	42 0
tgcacacgcc ggcgcctttc cggacaaatc gggatggcgt tccccaaaaa acagcttgat	48 0
gctgcaattt gttatccg gaaacacccc gaaatccgcg attgttaat ttcaggcggt	54 0
gatggctgc tcatcaacga ccaaattttt gaatatattt taaaagagcc ggcgcacact	60 0
ccgcacatccg aagtcatccg catcggaaaca cgtgctcccg tcgtcttcc gcagcgcatt	66 0
accgatcatc tgtgcgagat attgaaaaaaa tatcatccgg tctggctgaa caccatttt	72 0
aacacaagca tcgaaatgac agaaatcc gttgaggcat gtgaaaagct ggtgaacgcg	78 0
ggagtgccgg tcggaaatca ggctgtgta ttagcaggta ttaatgattc ggttccaatt	84 0
gtgaaaaagc tcatgcata cttggtaaaa atcagagtcc gtccttattatattaccaa	90 0
tgtgatctgt cagaaggaat aaggcattcc cgtgctccgt tttccaaagg ttggagatc	96 0
attgaagggc tgagaggtca tacccataggc tatgcggttc ctaccttgcgtt cgttacgc	102 0
ccaggcggag gaggtaaaat cgccatgcag ccgaactatg tcctgtctca aagtcctgac	108 0
aaagtgtatc taagaaattt tgaaagggtgt attacgtcat atccggaaacc agagaattat	114 0
atccccaaatc aggcagacgc ctattttgcgttcc tccgtttcc ctgaaaccgc tgacaaaaag	120 0
gagccgatcg ggctgagtgc cattttgct gacaaagaag tttcgtctac acctgaaaat	126 0
gtagacagaa tcaaacggcg tgaggcatac atcgcaatc cggagcatga aacattaaaa	132 0
gatcggcgtg agaaaagagg tcagctcaaa gaaaagaaat tttggcgca gcagaaaaaa	138 0
cagaaagaga ctgaatgcgg aggggattct tcataa	141 6

&lt;210&gt; 16

&lt;211&gt; 471

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<212> PRT  
<213> Artificial Sequence

<220>  
<223> Synthetic Construct

<400> 16

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Glu Glu Ile Glu  
1 5 10 15

Arg Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln  
20 25 30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Pro Arg Ser Thr Pro His Leu Glu Val Ile Arg Ile  
195 200 205

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Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Val Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Ser Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

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Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
 435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
 450 455 460

Glu Cys Gly Gly Asp Ser Ser  
 465 470

<210> 17  
 <211> 1416  
 <212> DNA  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<400> 17		
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gatgattt	aa agaaaagtcat taatctgacc gaggatgaag aggaaggcgt ccgtatttct	180
accaaaacg	a tccccttaaa tattacacct tactatgctc cttaatgga ccccgacaat	240
ccgagatg	cc cggtacgcat gcagtctgtg ccgcattccg aagaaaatgca caaaacaaaa	300
tacgata	tg aagacccgct tcatgaggat gaagatacac cggtacccgg tccgacaacac	360
cgctatcc	cg accgtgtgct gtttcttgta acgaatcaat gctccgtgta ctgcgcac	420
tgcacacg	cc ggcacaccc cgacaaatc ggaatggcg tccccaaaaa acagcttgc	480
gctgcaat	tg cttatatccg gaaaacaccc gaaatcccg atttttaat ttcaaggcggt	540
gatgggct	gc tcatcaacga ccaaatttttta gaatatattt taaaagagct ggcgcacatt	600
ccgcac	tc aagtcatccg catcggaaaca cgtgctccg tcgtcttcc gcagcgcatt	660
accgatcat	tc tgtgcgagat attgaaaaaa tatcatccgg tctggctgaa cacccatttt	720
aacacaag	ca tcgaaaatgac agaagaatcc gttgaggcat gtgaaaagct ggtgaacg	780
ggagtgc	cc ggaaatca ggctgtcgta ttagcaggta ttaatgattt ggttccatt	840
atgaaaaa	gc tcatgcata cttggtaaaa atcagagttcc gtccttatta tatttaccaa	900
tgtgatct	gt cagaaggaat aaggcatttc cgtgctccg tttccaaagg tttggagatc	960
attgaagg	tc tgagaggtca tacctcaggc tatgcgggtc ctaccttgcgatc gttcac	1020
ccaggcgg	gc gaggtaaaat ccgcctgcag ccgaactatg tcctgtctca aagtccctgac	1080
aaagtgat	tc taagaaattt tgaaggtgtg attacgtcat atccggaaacc agagaattat	1140

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atccccaaatc	aggcagacgc	ctatttgag	tccgtttcc	ctgaaaccgc	tgacaaaaaag	1200
gagccgatcg	ggctgagtgc	cattttgct	gacaaagaag	tttcgtctac	acctgaaaaat	1260
gtagacagaa	tcaaacggcg	tgaggcatac	atcgcaaatc	cggagcatga	aacattaaaa	1320
gatcggcgtg	agaaaagagg	tcagctaaa	gaaaagaaaat	ttttggcgca	gcagaaaaaa	1380
cagaaagaga	ctgaatgcgg	aggggattct	tcataa			1416

<210> 18  
 <211> 471  
 <212> PRT  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<400> 18

Met	Lys	Asn	Lys	Trp	Tyr	Lys	Pro	Lys	Arg	His	Trp	Lys	Glu	Ile	Glu
1				5				10					15		

Leu	Trp	Lys	Asp	Val	Pro	Glu	Glu	Lys	Trp	Asn	Asp	Trp	Leu	Trp	Arg
				20				25					30		

Leu	Thr	His	Thr	Val	Arg	Thr	Leu	Asp	Asp	Leu	Lys	Lys	Val	Ile	Asn
				35				40					45		

Leu	Thr	Glu	Asp	Glu	Glu	Gly	Val	Arg	Ile	Ser	Thr	Lys	Thr	Ile	
				50			55					60			

Pro	Leu	Asn	Ile	Thr	Pro	Tyr	Tyr	Ala	Pro	Leu	Met	Asp	Pro	Asp	Asn
65					70				75				80		

Pro	Arg	Cys	Pro	Val	Arg	Met	Gln	Ser	Val	Pro	Leu	Ser	Glu	Glu	Met
				85				90					95		

His	Lys	Thr	Lys	Tyr	Asp	Met	Glu	Asp	Pro	Leu	His	Glu	Asp	Glu	Asp
					100			105					110		

Thr	Pro	Val	Pro	Gly	Pro	Thr	His	Arg	Tyr	Pro	Asp	Arg	Val	Leu	Phe
					115			120				125			

Leu	Val	Thr	Asn	Gln	Cys	Ser	Val	Tyr	Cys	Arg	His	Cys	Thr	Arg	Arg
					130			135				140			

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Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

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Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
 385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
 405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
 420 425 430

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
 450 455 460

Glu Cys Gly Gly Asp Ser Ser  
465 470

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aacacaagca	tcgaaatgac	agaagaatcc	gttgaggcat	gtaaaaagct	ggtgaac <del>gc</del> cg	780
ggagtgc <del>cc</del> gg	tcggaaatca	ggctgtcgta	ttagcaggta	ttaatgattc	ggttccaa <del>t</del> tt	840
atgaaaaagc	tcatgc <del>at</del> ga	cttggtaaaa	atcagagtcc	gtc <del>c</del> tttatta	tat <del>t</del> acc <del>aa</del>	900
tgtatctgt	ctgagggctt	ggggcatttc	cgtgctcctg	tttccaaagg	tttggag <del>a</del> tc	960
attgaagggc	tgagaggtca	tacctcaggc	tatgcgg <del>tt</del> tc	ctac <del>c</del> ttgt	cg <del>t</del> tcac <del>gc</del> ca	1020
ccaggcggag	gaggtaaaat	cgc <del>c</del> ctgcag	ccgaactatg	tcctgtcaca	aagtcc <del>t</del> gac	1080
aaagtatct	taagaaat <del>ttt</del>	tgaagg <del>t</del> gtg	attacgtcat	atccggaacc	agagaatt <del>at</del>	1140
atccccaaatc	aggcagacgc	ctat <del>t</del> tgag	tccgttttcc	ctgaaaaccgc	tgacaaa <del>aa</del> ag	1200
gagccgatcg	ggctgagtgc	cattttgct	gacaaagaag	tttgc <del>t</del> ttac	ac <del>c</del> tgaaaat	1260
gt <del>a</del> gacagaa	tcaa <del>a</del> cggcg	tgaggcatac	atcg <del>c</del> aaatc	cggagcatga	aacatta <del>aa</del> aa	1320
gatcggcgtg	agaaaagaga	tcagctaaa	gaaaagaaaat	tttggcgca	gcagaaa <del>aa</del> aa	1380
cagaaagaga	ctgaatgc <del>gg</del>	agggattct	tcataa			1416

&lt;210&gt; 20

&lt;211&gt; 471

&lt;212&gt; PRT

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; Synthetic Construct

&lt;400&gt; 20

Met	Glu	Asn	Lys	Trp	Tyr	Lys	Pro	Lys	Arg	His	Trp	Lys	Glu	Ile	Glu
1							5		10					15	

Leu	Trp	Lys	Asp	Val	Pro	Glu	Glu	Lys	Trp	Asn	Asp	Trp	Leu	Trp	Gln
								20					25		30

Leu	Thr	His	Thr	Val	Arg	Thr	Leu	Asp	Asp	Leu	Lys	Lys	Val	Ile	Asn
								35					40		45

Leu	Thr	Glu	Asp	Glu	Glu	Gly	Val	Arg	Ile	Ser	Thr	Lys	Thr	Ile	
							50					55		60	

Pro	Leu	Asn	Ile	Thr	Pro	Tyr	Tyr	Ala	Ser	Leu	Met	Asp	Pro	Asp	Asn
							65				70		75		80

Pro	Arg	Cys	Pro	Val	Arg	Met	Gln	Ser	Val	Pro	Leu	Ser	Glu	Glu	Met
							85				90			95	

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His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Leu Gly His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

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Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
 325 330 335

Val Val His Ala Pro Gly Gly Gly Lys Ile Ala Leu Gln Pro Asn  
 340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
 355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
 370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
 385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Phe  
 405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
 420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Asp Gln  
 435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
 450 455 460

Glu Cys Gly Gly Asp Ser Ser  
 465 470

<210> 21  
 <211> 1416  
 <212> DNA  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<400> 21  
 atgaaaaaca aatggtataa accgaaacgg cattggaagg agatcgagt t atggaaggac 60  
 gtcccgaaag agaaatggaa cgattggctt tgacagctga cacacactgt aagaacgtta 120  
 gatgatttaa agaaagtcat taatctgacc gaggatgagg aggaaggcgt ccgtatttct 180  
 accaaaaacga tccccttaaa tattacacct taccatgctt cttaatgga ccccgacaat 240  
 ccgagatgcc cggtacgcat gcagtctgtg ccgcattctg aagaaatgca caaaacaaaa 300

tacgacatgg aagacccgct tcatgaggat gaagattcac cggtaaccgg tccgacacac	360
cgctatcccg accgtgtgct gtttcttgc acgaatcaat gttccgtgta ctgccgccac	420
tgcacacgccc ggctctttc cggacaaaatc ggaatggcg tccccaaaaa acagcttgat	480
gctgcaattg cttatatccg ggaaacacacc gaaatccgat attgttaat tt <del>c</del> aggcggt	540
gatgggctgc tcatcaacga ccaaatttttta gaatatattt taaaagagct g <del>c</del> gcagcatt	600
ccgcacatctgg aagtcatccg catcgaaaca cgtgctcccg tcgtcttcc gc <del>a</del> gcgcgtt	660
accgatcatc tgtgcgagat attgaaaaaa tatcatccgg tctggctgaa c <del>ac</del> ccatctt	720
aacacaagca tcgaaatgac agaagaaccc gttgaggcat gtgaaaagct gg <del>t</del> gaacgcg	780
ggagtgccgg tcgaaatca ggctgtcgta ttagcgggta ttaatgattc gg <del>t</del> tccaatt	840
atgaaaaagc tcatgcatga cttggtaaaa atcagagtcc gtccttatta t <del>at</del> ttaccaa	900
tgtgatctgt cagaaggaat aaggcatttc tgtgctctg tttccaaagg t <del>t</del> tgagatc	960
attgaagggc tgagaggtca tacctcaggc tatcggttc ctaccttgcgta c <del>g</del> ttcacgca	1020
ccaggcggag gaggtaaaat cgccctgcag ccgaactatg tcctgtctca a <del>ag</del> tcctgac	1080
aaagtgatct taagaaattt tgaaggtgtg attacgtcat atccggagcc ag <del>a</del> gaattat	1140
atccccaaatc aggacacgc ctatttgag tccgtttcc ctgaaaccgc tgacaaaaag	1200
gagccgatcg ggctgagtgc cattttgct gacaaagaag tttcgctac ac <del>c</del> tgaaaat	1260
gtagacagaa tcaaacggcg tgaggcatac atcgcaaatc cggagcatga aacattaaaa	1320
gatcggcgtg agaaaagagg tcagctaaa gaaaagaaaat tttggcgca gc <del>a</del> aaaaaaa	1380
cagaaagaga ctgaatgcgg agggattct tcataa	1416

<210> 22  
 <211> 471  
 <212> PRT  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<400> 22

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu			
1	5	10	15

Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln		
20	25	30

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Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Asn Ile Thr Pro Tyr His Ala Ser Leu Met Asp Pro Asp Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Pro Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Leu Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Val Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Leu  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Pro Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

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Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Cys Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
450 455 460

Glu Cys Gly Gly Asp Ser Ser  
465 470

<210> 23  
<211> 1416  
<212> DNA  
<213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<400> 23  
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 gttccggacg aaaagtggaa cgattggctt tgacagctga cacacactgt aagaacgtta  
 gatgattcaa agaaagtcat taatctgacc gaggatgaag aggaaggcgt ccgtatttct 120  
 accaaaaacga tccccttaaa tattacacct tactatgctt cttaatgga ccccgacaaat 180  
 ccgagatgcc cggtacgcat gcagtctgtg ccactttctg aagaaatgca caaaacaaaa 240  
 tacgatatgg aagacccgct tcatgaggat gaagattcac cggtacccgg tctgacacac 300  
 cgctatcccg gccgtgtgct gtttcttgc acgaatcaat gttccgtgca ctgccgccac 360  
 tgcacacgccc ggcgcttttc cggacaaatc ggaatggcg tcccgaaaaa acagcttgc 420  
 gctgcaatttgc ttatatccg gaaaacaccc gaaatccgat attgttaat ttcaggcggt 480  
 gatgggctgc tcatcaacga ccaaatttta gaatataattt taaaagagct ggcgcgcatt 540  
 ccgcacatctgg aagtcatccg catcggaaaca cgtgctcccg tcgtctttcc gcagcgcatt 600  
 accgatcatc tgtgcgagat attgaaaaaa tatcatccgg tctggctgaa cacccatttt 660  
 aacacaagca tcgaaatgac agaagaatcc gttgaggcat gtgaaaagct ggtgaacgcg 720  
 ggagtgcgg tcgaaatca ggctgtcgta ttagcaggta ttaatgatcc ggttccaatt 780  
 atgaaaaagc tcatgcatga ttggtaaaa atcagagtcc gtccttatttatttaccaa 840  
 tgtgatctgt cagaaggaat aaggcatttc cgtgctcctg tttccaaagg tttggagatc 900  
 attgaaggc tgagaggtca tacctcaggc tatgcggttc ctacctttgt cgttcacgca 960  
 ccaggcggag gagtaaaaat cgccctgcag ccgaactatg tcctgtctca aagtcctgac 1020  
 aaagtgtatct taagaaattt tgaaggtgtg attacgtcat atccggaaacc agagaattat 1080  
 atccccaaatc aggcaacgcgc ctatttgag tccgttttcc ctgaaaccgc tgacaaaaag 1140  
 gagccgatcg ggctgagtgc cattttgtc ggcaaagaag tttcgatc acctgaaaat 1260  
 gtagacagaa tcaaacggcg tgaggcatac atcgcaaatc cggagcatga aacattaaaa 1320  
 gatcggcgtg agaaaagagg tcagctaaaa gaaaagaaaat tttggcgca gcagaaaaaa 1380  
 cagaaagaga ctgaatgcgg aggggattct tcataa 1416

<210> 24  
 <211> 471  
 <212> PRT  
 <213> Artificial Sequence

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&lt;220&gt;

&lt;223&gt; Synthetic Construct

&lt;400&gt; 24

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu  
1 5 10 15

Leu Trp Lys Asp Val Pro Asp Glu Lys Trp Asn Asp Trp Leu Trp Gln  
20 25 30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Ser Lys Lys Val Ile Asn  
35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Gly Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val His Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Glu Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

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Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Gly Lys Glu Val Ser Ser  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
435 440 445

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Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys **Glu** Thr  
 450 455 460

Glu Cys Gly Gly Asp Ser Ser  
 465 470

<210> 25  
 <211> 1416  
 <212> DNA  
 <213> Artificial Sequence  
  
 <220>  
 <223> Synthetic Construct

<400> 25  
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 gttccggaag agaaatggaa cgattggctt tgacagctga cacacactgt aagraacgttg 120  
 gatgatttaa agaaagtcat taacctgacc gaggatgaag aggaaggcgt ccgtatttct 180  
 accaaaaacga tccctttaaa tattacacct tactatgctt cttaatgga ccccgacaaa 240  
 ccgagatgcc cggtacgcat gcagtctgtg ccgccttctg aagaaatgca caaaacaaaa 300  
 tacgatatgg aagacccgct tcatgaggat gaagattcac cggtacccgg tctgacacac 360  
 cgctatcccg accgtgtgct gtttcttgc acgaatcaat gttccgtgta ctgcccac 420  
 tgcacacgccc ggccgttttc cggacaaaatc ggaatggcg tccccaaaaa acagcttgat 480  
 gctgcaattt cttatatccg ggaaacacccc gaaatcccg attgtttat ttcaggcggt 540  
 gatgggctgc tcatcaacga ccaaatttttta gaatataattt taaaagagct gcgcagcatt 600  
 ccgcacatctgg aagtcatccg catcgaaaca cgtgctcccg tcgtcttcc gcaagcgcatt 660  
 accgatcatc tgtgcgagat attgaaaaaa tatcatccgg tctggctgaa cacccatttt 720  
 aacacaagca tcgaaatgac agaagaatcc gttgaggcat gtgaaaagct ggtgaacgcg 780  
 ggagtgcgg tcggaaatca ggctgtcgta ttagcaggta ttaatgattc ggttccaatt 840  
 atgaaaaagc tcatgcatga cttggtaaaa atcagagtcc gtccttatta tatttaccaa 900  
 tgtgacctgt cagaaggaat aaggcatttc cgtgctcccg ttccaaagg ttgggatc 960  
 attgaaggc tgggaggtca tacctcaggc tatgcggttc ctaccttgcgttcacgca 1020  
 ccaggcggag gaggtaaaat cgccctgcgg ccgaactatg tcctgtctca aagtccctgac 1080  
 aaagtgtatct taagaaattt tgaaggtgtg attacgtcat atccggaaacc agagaattat 1140  
 atccccaaatc aggccagacgc ctatccatgg tccgtttcc ctgaaaccgc tgaacaagaag 1200

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gagccgatcg ggctgagtgc cattttgct gacaaagaag tttcgctac acctgaaaat 1260  
 gtagacagaa tcaaacggcg tgaggcatac atcgcaaatc cggagcatga aacattaaaa 1320  
 gatcggcgtg agaaaagagg tcagctcaa gaaaagaaaat tttggcgca gcagaaaaaa 1380  
 cagaaagaga ctgaatgcgg aggggattct tcataa 1416

<210> 26  
 <211> 471  
 <212> PRT  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct  
 <400> 26

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu  
 1 5 10 15

Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln  
 20 25 30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
 35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
 50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Lys  
 65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
 85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
 100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
 115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
 130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
 145 150 155 160

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Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Gly Ile  
305 310 315 320

Ile Glu Gly Leu Gly Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Gly Lys Ile Ala Leu Arg Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

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Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
 385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
 405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
 420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
 435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
 450 455 460

Glu Cys Gly Gly Asp Ser Ser  
 465 470

<210> 27

<211> 1416

<212> DNA

<213> Artificial Sequence

<220>

<223> Synthetic Construct

<400> 27

atgaaaaaca aatggtataa accgaaacgg	cattggaagg agatcgagtt atggaaggac	60
gttccgggag agaaatggaa cgattggctt	tgacagctga cacacactgt aagaacgtta	120
gatgatttaa agaaaagtcat taatctgacc	gaggatgaag aggaaggcgt ccgtatttct	180
accaaaacga tccccttaaa tattacacct	tgctatgctc cttaatgga ccccgacaac	240
ccgagatgcc cggtacgcat gcagttgtg	ccgctttctg aagaaatgca caaaacaaaa	300
tacgatatgg aagacccgct tcgtgaggat	gaagattcac cggtacccgg tctgacacac	360
cgctatcccg accgtgtgct gtttcttgc	acgaatcaat gttccgtgta ctgccgccac	420
tgcacacgcc ggcgttttc cggacaaatc	ggaatggcgc tccccaaaaa acagcttgat	480
gctgcaattg cttatatccg ggaaacaccc	. gaaatccgcg attgtttaat ttcaggcgg	540
gatgggctgc tcatcaacgg ccaaatttta	gaatataattt taaaagagct ggcgcac	600
ccgcacatctgg aagtcatccg catcgaaaca	cgtgctcccg tcgtcttcc gcagcgcatt	660
accgatcatc tgtgcgagat attgaaaaaa	tatcatccgg tctggctgaa cacccattt	720
aacacaagcg tcgaaatgac agaagaatcc	gttgaggcat gtgaaaagct ggtgaacgcg	780

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ggagtgcgg	tcggaaatca	ggctgtcgta	ttagcaggtta	ttaatgattc	ggttccaatt	840
atgaaaaagc	tcatgcatga	cttggtaaaa	atcagagtcc	gtccttatta	tatcaccaa	900
tgtgatctgt	cagaaggaat	aaggcatttc	cgtgctcctg	tttccaaagg	tttggagatc	960
attgaagggc	tgagaggtca	tacctcaggc	tatgcggttc	ctaccttgc	cgttacgca	1020
ccaggcggag	ggggtaaaat	cgcctgcag	ccgaactatg	tcctgtctca	aagtccctgac	1080
aaagtaatct	taagaaattt	tgaagggtgt	attacgtcat	atccggaacc	agagaattat	1140
atccccaaatc	aggcagacgc	ctattttgag	tccgtttcc	ctggaaccgc	tgacaaaaag	1200
gagccgatcg	ggctgagtgc	cattttgct	gacaaagaag	tttcgtctac	acctgaaaat	1260
gtagacagaa	tcaaacggcg	tgaggcatac	atcgcaaatc	cggagcatga	aacattaaaa	1320
gatcggcgtg	agaaaagagg	tcagctaaa	gaaaagaaat	cttggcgca	gcagaaaaaa	1380
cagaaagaga	ctgaatgcgg	aggggattct	tcataa			1416

&lt;210&gt; 28

&lt;211&gt; 471

&lt;212&gt; PRT

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; Synthetic Construct

&lt;400&gt; 28

Met	Lys	Asn	Lys	Trp	Tyr	Lys	Pro	Lys	Arg	His	Trp	Lys	Glu	Ile	Glu
1				5				10					15		

Leu	Trp	Lys	Asp	Val	Pro	Gly	Glu	Lys	Trp	Asn	Asp	Trp	Leu	Trp	Gln
				20				25					30		

Leu	Thr	His	Thr	Val	Arg	Thr	Leu	Asp	Asp	Leu	Lys	Lys	Val	Ile	Asn
				35				40					45		

Leu	Thr	Glu	Asp	Glu	Glu	Gly	Val	Arg	Ile	Ser	Thr	Lys	Thr	Ile	
				50			55		60						

Pro	Leu	Asn	Ile	Thr	Pro	Cys	Tyr	Ala	Pro	Leu	Met	Asp	Pro	Asp	Asn
65					70				75				80		

Pro	Arg	Cys	Pro	Val	Arg	Met	Gln	Ser	Val	Pro	Leu	Ser	Glu	Glu	Met
				85				90					95		

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His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu Arg Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Gly Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Val Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

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Val Val His Ala Pro Gly Gly Gly Lys Ile Ala Leu Gln Pro Asn  
 340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
 355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
 370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Gly Thr Ala Asp Lys Lys  
 385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
 405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
 420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
 435 440 445

Leu Lys Glu Lys Lys Ser Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
 450 455 460

Glu Cys Gly Gly Asp Ser Ser  
 465 470

<210> 29

<211> 1416

<212> DNA

<213> Artificial Sequence

<220>

<223> Synthetic Construct

<400> 29

atgaaaaaaca aatggtataa accgaaacgg cattggaagg agatcgagtt atggaaggac 60

gttccggaag agaaatggaa cgattggctt tgacggctga cacacactgt aagaacgtta 120

gatgatttaa agaaagtcat taatctgacc gaggatgaag aggaaggcgt ccgtatttct 180

acccaaaacga tccccttaag tattacacct tactatgctt cttaatgga ccccgacaat 240

ccgagatgcc cggtacgcat gcagtctgtg ccgccttctg aggaaatgca caaaaacaaaa 300

tacgatatgg aagaccgcgt tcatgaggat gaagattcac cggtacccgg tctgacacac 360

cgctatcccc accgtgtgct gtttcttgc acgaatcaat gttccgtgta	ctgcccgcgc	420		
tgcacacgccc ggccgttttc cggacagatc ggaatggcg tccccaaaaa	acagcttgat	480		
gctgcaattg cttatatccg ggaaacaccc gaaatccgat	attgttaat	ttcaggcggt	540	
gatgggctgc tcatcaacga ccaaattta gaatataattt	taaaagagct	gcgcagcatt	600	
ccgcacatctgg aagtcatccg catcggaca cgtgctcccg	tcgtcttcc	gcagcgcatt	660	
accgatcatc tgtgcgagat attgaaaaaa tatcatccgg	tctggctgaa	cacccatttt	720	
aacacaagca tcgaaatgac agaagaatcc gttgaggcat	gtgaaaagct	gtgaacgcg	780	
ggagtgccgg tcggaaatca ggctgtcgta ttagcaggtt	ttaatgattc	ggttccaatt	840	
atgaaaaaagc tcatgcatga cctggtaaaa atcagagtcc	gtccttatta	tatccatcaa	900	
tgtgatctgt cagaaggaat acggcatttc cgtgctcctg	tttccaaagg	tttggagatc	960	
attgaagggc tgagaggtca tacctcaggc tatgcggttc	ctacctttgt	gttcacgca	1020	
ccaggcggag gaggtaaaat cgcctgcag ccgaactatg	tcctgtctca	agtcctgac	1080	
aaagtatct taagaaattt tgaagggtgt	attacgtcat	atccggaaacc	agagaattat	1140
atccccaaatc aggcagacgc ctatttgag tccgtttcc	ctgaaaccgc	tgcacaaaaag	1200	
gagccgatcg ggctgagtgc cattttgct	gacaaagaag	ttcgtctac	acctgaaaat	1260
gtagacagaa tcaaacggcg tgaggcatac atcgcaaatc	cgagcatga	macattaaaa	1320	
gatcggcgtg agaaaagagg tcagctaaa gaaaagaaaat	ttttggcgca	gcagaaaaaa	1380	
cagaaagaga ctgaatgcgg agggattct tcataa			1416	

<210> 30  
 <211> 471  
 <212> PRT  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<400> 30

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu			
1	5	10	15

Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Arg		
20	25	30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn		
35	40	45

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Leu Thr Glu Asp Glu Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Ser Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg Arg Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

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Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
450 455 460

Glu Cys Gly Gly Asp Ser Ser  
465 470

<210> 31  
<211> 1416  
<212> DNA  
<213> Artificial Sequence

<220>

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&lt;223&gt; Synthetic Construct

<400> 31	
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gttccggaag agaaatggaa cgattggctt tgacagctga cacgcactgt aagaacgtta	120
gatgattaa agaaagtcat taatctgacc gaggatgaag aggaaggcgt ccgtatttct	180
accaaaaacga tccccttaaa tattacacct tactatgcga gcttaatgga tccagaaac	240
ccacgttgc cggtacgcat gcagtctgtg ccgccttctg aagaaatgca cacaagcaaa	300
tatgacatgg aagatccgct tcatgaggat gaagattcac cggtacccgg tctgacacac	360
cgctatcccg accgtgtgct gtttcttgc acgagtcaat gtcccgta ctgccgccac	420
tgcacacgcc ggcgcctttc cggacaaatc ggaatggcg tccccaaaaa acagcttgat	480
gctgcaattg cttatatccg gcaaacaccc gaaatcccg attgttaat ttcaggcgt	540
gatgggctgc tcatcaacga ccaaattta gaatatattt taaaagagct ggcgcaggatt	600
ccgcacatctgg gagtcatccg catcggaaca cgtgctcccg tcgtcttcc gcagcgcatt	660
accgatcatc tgtgcgagat attgaaaaga tatcatccgg tctggctgaa cacccatttt	720
aacacaagca tcgaaatgac agaagaatcc gttgaggcat gtgaaaagct ggtgaacgac	780
ggagtgccgg tcggaaatca ggctgtcgta ttagcaggta ttaatgattc gtttccaaatt	840
atgaaaaagc tcatgcatga cttggtaaaa atcagagtcc gtccttatta tatttaccaa	900
tgtgatctgt cagaaggaat aaggcatttc cgtgctcctg tttccaaagg tttggagatc	960
attgaagggc tgagaggtca tacctcaggc tatgcggttc ctacccttgc cgttcacgca	1020
ccaggcggag gaggtaaaat cgcctgcag ccgaactatg tcctgtctca aagtccgtac	1080
aaagtgtatc taagaaattt tgaagggtgt attacgtcat atccgaaacc agagaattat	1140
atccccaaatc aggcagacgc ctatttgag tccgtttcc ctgaaaccgc tgacaaaaag	1200
gagccgatcg ggctgagtgc cattttgct gacaaagaag tttcgctac acctgaaaat	1260
gtagacagaa tcaaacggcg tgaggcatac atcgaaatc cggagcatga aacattaaa	1320
gatcggcgtg agaaaagagg tcagctaaa gaaaagaaaat tttggcgca gcagaaaaaa	1380
cagaaaagaga ctgaatgcgg agggattct tcataa	1416

&lt;210&gt; 32

&lt;211&gt; 471

&lt;212&gt; PRT

&lt;213&gt; Artificial Sequence

&lt;220&gt;

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<223> Synthetic Construct

<400> 32

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu  
1 5 10 15

Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Glu  
20 25 30

Leu Thr Arg Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
35 40 45

Leu Thr Glu Asp Glu Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Glu Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Thr Ser Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Ser Gln Cys Pro Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Gly Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

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Cys Glu Ile Leu Lys Arg Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
 450 455 460

Glu Cys Gly Gly Asp Ser Ser  
 465 470

<210> 33  
 <211> 1416  
 <212> DNA  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<400> 33  
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 gttccggaag agaaatggaa cgattggctt tgacagctga cacac actgt aagaacgtta 120  
 gatgatttaa agaaagtcat taatctgacc gaggatgaag aggaa ggcgt ccgtatttct 180  
 accaaaaacga tccccttaaa tattacacct tactatgctt cttaatgga cccgacaat 240  
 ccgagatgcc cggtacgcat gcagtctgtg ccgcattctg aagaa atgca caaaacaaaa 300  
 tacgatatgg aagacccgct tcatgaggat gaagattcac cggtacccgg tctgacacac 360  
 cgctatcccg accgtgtgtct gttcttgtc acgaatcaat gttccgtgta ctgccgccac 420  
 tgcacacgccc ggcgcatttc cggacaaatc ggaatggcg tccccaaaaa acagcttgat 480  
 gctgcaatttgc ttatatccg ggaaacaccc gaaatccgca actgt ctgtt gtctggcggt 540  
 gatgggctgc tcatcaacga ccaaatttttta gaatataattt taaaagagct ggcgcacattt 600  
 ccgcattctgg aagtcatccg catcggaca cgtgctcccg tcgtcttcc gcagcgcattt 660  
 accgatcacc tgtgcgagat gttaaaaaaaaa tatcatccgg tctggctgaa cacccatttt 720  
 aacacaagca tcgaaatgac agaagaatcc gttgaggcat gtgaaaagct ggtgaacgcg 780  
 ggagtgcgcgg tcggaaatca ggctgtcgta ttagcaggta ttaatgatttgc gttccaattt 840  
 atgaaaaaagc tcatgcatga ctggtaaaa atcagagtcc gtccttatttta tatttaccaa 900  
 tgtgatctgt cagaaggaat aaggcatttc cgtgctccgt tttccaaagg tttggagatc 960  
 attgaagggc tgagaggtca tacctcaggc tatgcggttc ctacc tttgt cgttcacgca 1020  
 ccaggcggag gaggtaaaat cgccctgcag ccgaactatg tcctgtctca aagtcctgac 1080  
 aaagtgatct taagaaattt tgaaggtgtg attacgtcat atccggaaacc agagaattat 1140  
 atcccccaatc aggcagacgc ctattttagt cccgtttcc ctgaaaccgc tgacaaaaag 1200  
 gagccgatcg ggctgagtgc gctgtttgct gacaaagaag tttcg tctac acctgaaaat 1260

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gtagacagaa tcaaacggcg tgaggcatac atcgcaaatc cggagcatga aacattaaaa 1320

gatcggcgtg agaaaagagg tcagctaaa gaaaagaaaat ttttggcgca gcagaaaaaa 1380

cagaaagaga ctgaatgcgg aggggattct tcataa 1416

<210> 34

<211> 471

<212> PRT

<213> Artificial Sequence

<220>

<223> Synthetic Construct

<400> 34

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu  
1 5 10 15

Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln  
20 25 30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

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Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Leu Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Met Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Leu Phe Ala Asp Lys Glu Val Ser Ser  
 405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
 420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
 435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
 450 455 460

Glu Cys Gly Gly Asp Ser Ser  
 465 470

<210> 35

<211> 1416

<212> DNA

<213> Artificial Sequence

<220>

<223> Synthetic Construct

<400> 35

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gttccggaag agaaatggaa cgattggctt tgacagctg a cacacactgt aagaacgtta 120

gatgatttaa agaaagtcat taatctgacc gaggatgaag aggaaggcgt ccgtatttct 180

accaaaacga tccccttaaa ttcacacact tactatgcg a cttaatggc tccagaaaaac 240

ccacgttgc cggtaacgcat gcagtctgtg ccgtttcctg aagaaatgca caaaacaaaa 300

tacgatatgg aagacccgct tcatgaggat gaagattcac cggtaacccgg tctgacacac 360

cgctatcccg accgtgtgct gtttcttgc acggatcaat gttccgtgta ctgccgccac 420

cgcacacgcc ggcgttctc cggacaaatc ggaatggc tccccaaaa acagctgtat 480

gctgcaattt gtttccatccg ggaaacacccc gaaatccgc attgtttat ttcaggcggt 540

gatgggctgc tcatcaacga ccaaattttt gaatatattt taaaagagct ggcgcgcatt 600

ccgcattctgg aagtcatccg catcgaaaca cgtgtcccg tcgtttcc gcagcgcatt 660

accgatcatc tgtgcgagat attgaaaaaa catcatccgg tctggctgaa caccatttt 720

aacacaagca tcgaaatgac agaagaatcc gttgaggcat atgaaaagct ggtgaacgcg 780

ggagtgcggg tcggaaatca ggctgttgta ttagcaggtt ttaatgattc ggttccaatt 840

ataaaaaaagc tcatgcata	cttggtaaaa atcagagtcc	gtcccttatta tatttaccaa	900
tgtgacctgt cagaaggaat aaggcatttc	cgtgctcctg ttccaaagg	tttggagatc	960
attgaagggc tgagaggtca tacctcaggc	tatgcgggtc ctacctttgt	cgttcacgca	1020
ccaggcggag gaggtaaaat cgcctgcag	ccgaactatg tccctgtctca	aagtcctgac	1080
aaagtgtatc taagaaattt tgaaggtgtg	attacgtcat atccggaacc	agagaattat	1140
atccccaaatc aggagacgc ctatttgag	tccgttttcc ctgaaaccgc	tgacaaaaag	1200
gagccgatcg ggctgagtgc cattttgct	gacaaagaag tttcgtctac	acctgaaaat	1260
gtagacagaa tcaaacggcg tgaggcatac	atgccaaatc cggagcatga	aacattaaaa	1320
gatcggcgtg agaaaagagg tcagctaaa	gaaaagaaaat ttttggcgca	gcagaaaaaa	1380
cagaaagaga ctgaatgcgg	agggattct tcataa		1416

&lt;210&gt; 36

&lt;211&gt; 471

&lt;212&gt; PRT

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; Synthetic Construct

&lt;400&gt; 36

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu			
1	5	10	15

Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln			
20	25	30	

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn			
35	40	45	

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile			
50	55	60	

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Glu Asn			
65	70	75	80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Pro Glu Glu Met			
85	90	95	

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp			
100	105	110	

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Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asp Gln Cys Ser Val Tyr Cys Arg His Arg Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Glu Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys His His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Tyr Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Ile Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

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Val Val His Ala Pro Gly Gly Gly Lys Ile Ala Leu Gln Pro Asn  
 340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
 355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
 370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
 385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
 405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
 420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
 435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
 450 455 460

Glu Cys Gly Gly Asp Ser Ser  
 465 470

<210> 37  
 <211> 1416  
 <212> DNA  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<400> 37  
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 gatgatttaa agaaaagtcat taatctgacc gaggatgaag aggaaggcgt ccgtatttct 180  
 accaaaacga tccccttaaa tattacacct tactatgctt cttaatgga ccccgacaat 240  
 ccgagatgcc cggtacgcat gcagttgtg ccgtttctg aagaaaatgca caaaaa 300  
 tacgataatgg aagacccgct tcatgaggat gaagattcac cggtacccgg tctgacacac 360  
 cgctatccca accgtgtgct gtttcttgc acgaatcaat gttccgtgta ctgccgccac 420

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tgcacacgccc	ggcgctttc	cgacaaatc	gaaatggcg	tcccaaaaaa	acagcttgat	480	
gctgcaattg	cttataatccg	ggaaacaccc	gaaatcccg	actgtctgtt	gtctggcggt	540	
gatgggctgc	tcatcaacga	ccaaatttt	aatatattt	taaaagagct	gcgcagcatt	600	
ccgcatctgg	aagtcatcg	tatcggttct	cgtgcgcag	tcgtctttcc	gcagcgcatt	660	
accgatcatc	tgtgcgagat	attgaaaaaa	tatcatccgg	tctggctgaa	caccat	720	
aacacaagca	tcgaaatgac	agaagaatcc	gttggggcat	gtgaaaagct	ggtgaacg	780	
ggagtgc	tcggaaatca	ggctgtcgta	ttagcaggta	ttaatgattc	ggttccaatt	840	
atgaaaaagc	tcatgcatga	cttggtaaaa	atcagagtcc	gtccttatta	tat	900	
tgtgatctgt	cagaaggaat	agggcatttc	cgtgcctc	tttccaaagg	tttggagatc	960	
attgaagggc	tgagaggtca	tacctcaggc	tatcggttcc	ctacctt	gttca	1020	
ccaggcggag	gaggtaaaat	cgcctgcag	ccgaactatg	tcctgtcaca	aagtcc	1080	
aaagtgatct	taagaaattt	tgaaggtgt	attacgtcat	atccggaaacc	agagaattat	1140	
atccccaaatc	aggcagacgc	ctat	tttgag	tccgtttcc	tgaca	1200	
gagccgatcg	ggctgagtgc	cattttgct	gacaaagaag	tttcgtt	ac	1260	
gtagacagaa	tcaaacggcg	tgaggcatac	atgc	aaatc	cgagcat	1320	
gatcggcgtg	agaaaagaga	tca	gctcaaa	gaaaagaaat	tttggcgca	gcagaaaaaa	1380
cagaaagaga	ctgaatgcgg	aggggattct	tcataa			1416	

&lt;210&gt; 38

&lt;211&gt; 471

&lt;212&gt; PRT

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; Synthetic Construct

&lt;400&gt; 38

Met	Lys	Asn	Lys	Trp	Tyr	Lys	Pro	Lys	Arg	His	Trp	Lys	Glu	Ile	Glu
1							5			10			15		

Leu	Trp	Lys	Asp	Val	Pro	Glu	Glu	Lys	Trp	Asn	Asp	Trp	Leu	Trp	Gln
													20	25	30

Leu	Thr	His	Thr	Val	Arg	Thr	Leu	Asp	Asp	Leu	Lys	Lys	Val	Ile	Asn
													35	40	45

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Leu Thr Glu Asp Glu Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asn Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Leu Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Ser Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

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Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Gly His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Phe  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Asp Gln  
435 440 445

Leu Lys Glu Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
450 455 460

Glu Cys Gly Gly Asp Ser Ser  
465 470

<210> 39  
<211> 1416  
<212> DNA  
<213> Artificial Sequence

<220>  
<223> Synthetic Construct

<400> 39  
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 gatgatttaa agaaagtcat taatctgacc gaggatgaag aggaaggcgt ccgtatttct 180  
 accaaaacga tccccttaaa tattacacct tactatgctt ctttaatggc ccccgacaaat 240  
 ccgagatgcc cggtaacgcat gcagtcgtg ccgccttctg aagaaatgca caaaacaaaa 300  
 tacgatatgg aagacccgct tcatgaggat gaagattcac cggtaacccgg tctgacacac 360  
 cgctatcccg accgtgtgct gtttctgtc acgaatcaat gttccgtgca ctgcccac 420  
 tgcacacgcc ggcccttttc cggacaaatc ggaatggcg tccccaaaaa acagcttgat 480  
 gctgcaatttgc ttatatccg gaaaacaccc gaaatcccgat attgttaat ttcaggcggt 540  
 gatggctgc tcatcaacga ccaaatttttta gaatatatttt taaaagagct gcgcagcatt 600  
 ccgcacctgg aagtcatccg catcgaaaca cgtgctcccg tgcgtttcc gcagcgcatt 660  
 accgatcatc tgtgcgagat attgaaaaaa tatcatccgg tctggctgaa cacccatttt 720  
 aacacaagca tcgaaaatgac agaagaatcc gttgaggcat gtgaaaagct ggtgaacgca 780  
 ggagtgcgg tcggaaatca ggctgtcgta ttagcaggta ttaatgattt ggttccaatt 840  
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 tgtgatctgt cagaaggaat aaggcatttc cgtgctccgt tttccaaagg tttggagatc 960  
 attgaagggc tgagaggtca tacctcaggc tatgcgggttc ctaccttgcgt cgttcaac 1020  
 ccaggcggag gtggtaaaat cgcctgcag ccgaactatg tccctgtctca aagtcctgac 1080  
 aaagtgtatct taagaaattt tgaaggtgtg attacgtcat atccggaaacc agagaattat 1140  
 atccccaaatc aggccagacgc ctatttgag tccgtttcc ctgaaaccgc tgacaaaaag 1200  
 gagccgatcg ggctgagtc cattttgtc ggcaaaagaag tttcgtctac acctgaaaat 1260  
 gtagtcagaa tcaaacggcg tgaggcatac atcgcaaatc cggagcatga aacattaaaa 1320  
 gatcgccgtg agaaaagagg tcagctaaa gaaaagaaaat ttttggcgca gcagaaaaaa 1380  
 cagaaagaga ctgaatgcgg aggggattct tcataa 1416

<210> 40  
 <211> 471  
 <212> PRT  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

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<400> 40

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu  
1 5 10 15

Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln  
20 25 30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val His Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

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Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Gly Lys Glu Val Ser Ser  
405 410 415

Thr Pro Glu Asn Val Val Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
450 455 460

Glu Cys Gly Gly Asp Ser Ser  
 465 470

<210> 41  
<211> 1416  
<212> DNA  
<213> Artificial Sequence  
  
<220>  
<223> Synthetic Construct  
  
<400> 41  
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gtccccggaaag agaaaatggaa cgattggctt tgacagctga cacacactgt aagaacgtta  
gatgattaa agaaaagtcat taatctgacc gaggatgaaag aggaaggcgt ccgtatttct 120  
accaaaaacga tccccttaaa tattacacct tactatgctt cttaatgga ccccgacaat  
ccgagggtgcc cggtacgcat gcagtcgttg ccactgtctg agggaaatgca caaaagcaaa 180  
tatgacatgg aagatccgct tcatgaggat gaagattcac cggtacccgg tctgacacac  
cgctatcccg accgtgtgct gttcttgtc acgaatcaat gttccgtgta ctgcccgcac 240  
tgcacacgccc ggcgcttttc cggacaaatc ggaatgcgcg tccccaaaaaa acagcttgat  
gctgcaatttgc ttatatccg ggaaacacccc gaaatccgcg attgttaat ttcaggcggt 300  
gatgggctgc tcatcaacga ccaaatttttta gaatataattt taaaagagct ggcgcgcatt  
ccgcacatctgg aagtcatccg catcggaaaca cgtgctcccg tcgtcttcc gcagcgcatt 360  
accgatcatc tgtgcgagat attgaaaaaa tatcatccgg tctggctgaa caccatttt  
aacacaagca tcgaaatgac agaagaatcc gttgaggcat gtgaaaagct ggtgaacgcg 420  
ggagtgcggg tcggaaatca ggctgtcgta ttagcaggta ttaatgattc gggttccaaatt  
atgaaaaaagc tcatgcatttgc cttggtaaaa atcagagtcg gtccttatttatttaccaa 480  
tgtgatctgt cagaaggaat aaggcatttc cgtgctcccg tttccaaagg tttggagatc  
attgaagggc tgagaggta tacctcaggc tatgcgggttc ctacctttgt cggttccacgc 540  
ccggggcggag gaggtaaaaat cgcgcgtcag ccgaactatg tcctgtctca aagtccgtac  
aaagtgtatct taagaaattt tgaaggtgtg attacgtcat atccggaaacc agagaatttt  
atcccccaatc aggcagacgc ctatccgttgc tccgttttcc ctgaaaccgc tgacaaaaag 600  
gagccgatcg ggctgagtgc catttttgct gacaaaagaag tttcgtctac acctgaaaat  
gttagacagaa tcaaaccggcg tgaggcgtac atcgcaatc cgagacatga aacattaaaa 660  
1140  
1200  
1260  
1320

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gatcggcgtg agaaaagagg tcagctaaa gaaaagaaat ttttggcgca gcagaaaaaa 1380  
cagaaaagaga ctgaatgcgg aggggattct tcataa 1416

<210> 42  
<211> 471  
<212> PRT  
<213> Artificial Sequence

<220>  
<223> Synthetic Construct  
<400> 42

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu  
1 5 10 15

Leu Trp Arg Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln  
20 25 30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Ser Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

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Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

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Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
 405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
 420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
 435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
 450 455 460

Glu Cys Gly Gly Asp Ser Ser  
 465 470

&lt;210&gt; 43

&lt;211&gt; 1416

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; Synthetic Construct

<400> 43		
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gttccggaag agaaatggaa cgattggctt tgacagctga cacacactgt aagaacgtta	120	
gatgatttaa agaaaagtcat taatctgacc gaggatgaag aggaaggcgt ccgtatttct	180	
accaaaaacga tccccttaaa tattacacca tactatgcga gcttaatgga tccagaaaaac	240	
ccacgttgtc cgttacgcatt gcagttgttg ccgtttccg aagaaatgca caaaaacaaaa	300	
tacgatatgg aagacccgct tcatgaggat gaagattcac cggtacccgg tctgacacac	360	
cgctatcccg accgtgtgtc gtttcttgtc acgaatcaat gttccgtgttta ctgccgccac	420	
tgcacacgccc ggcgttttc cggacaaatc ggaatggcg tccccaaaaa acagcttgcgt	480	
gctgcaatttgc ttatatccg ggaaacacccc gaaatcccg attgtttat ttcagggcggt	540	
gatgggctgc tcatcaacga ccaaatttttta gaatataattt taaaagagct ggcgcgcatt	600	
ccgcattctgg aagtcatccg catcgaaaca cgtgctcccg tcgtcttcc gcagcgcatt	660	
accgatcatc cgtgcgagat attgaaaaaa tatcatccgg tctggctgaa cacccatttt	720	
aacacaagca tcgaaaatgac agaagaatcc gttgaggcat gtgaaaagct ggtgaacgcg	780	
ggagtgccgg tcggaaatca ggctgtcgta ttagcaggta ttaatgattc ggttccaatt	840	
atgaaaaaagc tcatgcatttgc cttggtaaaaa atcagagttcc gtccttattttatccaa	900	

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tgtgatctgt cagaaggaat aaggcattc cgtgctcctg tctccaaagg	960
tttggagatc attgaagggc tgagaggtca taccccaggc tatgcgggtc ctaccttgc	1020
cgttcacgca ccaggcggag gaggtaaaat cgccctgcag ccgaactatg tcctgtctc	1080
aagtgcataa aaggtgatct taagaaattt tgaaggtgtg attacgtcat atccggaacc	1140
agagaattat atccccaaatc aggacacgc ctatttgag tccgtttccc ctgaaaccgc	1200
tgacaaaaag gagccgatcg ggctgagtgc cattttgct gacaaagaag tttcgctac	1260
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aacattaaaa gatcggcgtg agaaaagagg tcagctaaa gaaaagaaat tttcgccgc	1380
gcagaaaaaa cagaaagaga ctgaatgcgg agggattct tcataa	1416

<210> 44  
 <211> 471  
 <212> PRT  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct  
 <400> 44

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu			
1	5	10	15

Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln		
20	25	30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Val Ile Asn		
35	40	45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile		
50	55	60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Glu Asn			
65	70	75	80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met		
85	90	95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp		
100	105	110

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Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Pro  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Pro Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

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Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
 355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
 370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Ser Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
435 440 445

Leu Lys Glu Lys Lys Phe Ser Ala Gln Gln Lys Lys Gln Lys Glu Thr  
 450 455 460

Glu Cys Gly Gly Asp Ser Ser  
465 470

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<210> 45
<211> 1416
<212> DNA
<213> Artificial Sequence
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<220>  
<223> Synthetic Construct

<400> 45

atggaaaaaca aatggtataa accgaaacgg cattggaagg agatcgagtt acggaaggac 60  
gttccggaag agaaatggaa cgattggctt tgacagctga cgcacactgt aagaacgtta 120  
gatgatttaa agaaagtcat taatctgacc gaggatgaag aggaaggcgt cctgtatttct 180  
accaaaaacga tcccccttaaa tattacacct tactatgcga gcttaattga tccagaaaaac 240  
ccacgttgtc cggtacgcat gcagtctgcg ccgctgtctg aagaaatgca caaaacaaaa 300  
tacgataatgg aagacccgct tcatgaggat gaagattcac cggtaacctggg tctgacacac 360  
cgctatcccg accgtgtgct gtttcttgct acgaatcaat gttccgtgta ctgccgcccac 420  
tgcacacgccc ggcgcttttc cggacaaatc ggaacggggcg tccccaaaaaa acagcttgat 480

gctgcaactg cttatatccg ggaaacacccc gaaatccgcg attgttaat tccaggc <del>gg</del> gt	540
gatgggctgc tcataaacga ccaaatttta ggatatattt taaaagagct ggcgcgcatt	600
ccgcacatctgg aagtcatccg catcgaaaca cgtgcccccg tcggcttcc gcagcgcatt	660
accgatcatc tgtgcgagat attgaaaaaa tatcatccgg tctggctgaa cacccatttt	720
aacacaagca tcgaaatgac agaagaatcc gttgaggcat gtgaaaagct ggtgaac <del>gc</del> g	780
ggagtgcggg tcggaaatca ggctgtcgta ttagcaggta ttaatgattc ggttccaatt	840
atgaaaaagc tcatacatga ctggtaaaaa atcagagtcc gtccttatta tatttaccaa	900
tgtgatctgt cagaaggaat aaggcatttc cgtgctcctg tttccaaagg tttggagatc	960
attgaagggc tgagaggtca tacctcaggc tatgcggttc ctaccttgcgatc	1020
ccaggcggag gaggtaaaat cgcctgcag ccgaactatg ccctgtctca aagtcctgac	1080
aaagtgatct taagaaattt tgaaggtgtg attacgtcat atccgaaacc agagaattat	1140
atccccaaatc aggcagacgc ctatttgag tccgtttcc ctgaaaccgc tgacaaaaag	1200
gagccgatcg ggctgagtgc cattttgct gacaaagaag tttcgctac acctgaaaat	1260
gtagacagaa tcaaacggcg tgaggcatac atcgcaaatc cggagcatga aacattaaaa	1320
gatcggcgtg agaaaagagg tcagctaaa gaaaagaaaat tttggcgca gcagaaaaaa	1380
cagaaagaga ctgaatgcgg agggattct tcataa	1416

&lt;210&gt; 46

&lt;211&gt; 471

&lt;212&gt; PRT

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; Synthetic Construct

&lt;400&gt; 46

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu				
1	5		10	15
	10	15		

Leu Arg Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln		
20	25	30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn		
35	40	45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile		
50	55	60

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Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Ile Asp Pro Glu Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Ala Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Thr Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Thr Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Pro Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Gly Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Gly Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

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Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Ala Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
450 455 460

Glu Cys Gly Gly Asp Ser Ser  
465 470

<210> 47  
<211> 1416  
<212> DNA  
<213> Artificial Sequence

<220>  
<223> Synthetic Construct

<400> 47  
atggaaaaca aatggataa accgaaacgg cattggaagg agatcgagtt atggaaggac 60

gttccggaag	agaaaatggaa	cgattggctt	tgacagctga	cacaca <del>ctgt</del>	aagaacgtta	120
gatgatttaa	agaaaagtcat	taatctgacc	gaggatgaag	aggaaggcgt	ccgtat <del>ttct</del>	180
accaaaaacga	tccccttaaa	tattacacct	tactatgcga	gcttaattga	tccagaaaac	240
ccacgttgtc	cggtacgcat	gcagtctgtg	ccgcttccg	aagaaa <del>tgca</del>	caaaaacaaaa	300
tacgatatgg	aagatccgct	tcatgaggat	gaagattcac	cggta <del>ccgg</del>	cctgacacac	360
cgctatcccg	accgtgtgct	gtttcttgc	gcgaatcaat	gttccgtgta	ctgccgccac	420
tgcacacgcc	ggcgctttc	cggacaaatc	ggaatggcg	tcccc <del>aaaaa</del>	acagcttgat	480
gctgcaattg	cttataatccg	ggaaacacccc	gaaatccgcg	attgtt <del>taat</del>	ttcaggcggt	540
gatgggctgc	tcatcaacga	ccaaattttt	gaatataattt	taaaagagct	gcgcagcatt	600
ccgcacatccgg	aagtcatccg	catcggaaca	cgtgcccccg	tcgtctttcc	gcagcgcatt	660
accgatcatc	tgtgcgagat	attgaaaaaa	tatcatccgg	tctgg <del>ctgaa</del>	cacccatttt	720
aacacaagca	tcgaaatgac	agaagaatcc	gttgaggcat	gtgaaa <del>agct</del>	ggtgaacgcg	780
ggagtgccgg	tcggaaatca	ggctgtcgta	ttagcaggta	ttaatgattc	ggttccaatt	840
atgaaaaagc	tcatgcatga	cttggtaaaa	atcagagtcc	gtcctt <del>atta</del>	tat <del>tt</del> accaa	900
tgtgatctgt	cagaaggaat	aaggcatttc	cgtgcccctg	tttcc <del>aa</del> agg	tttggagatc	960
attgaagggc	tgagaggtca	tacctcaggc	tgtgcggttc	ctac <del>ctt</del> gt	cgttcacgca	1020
ccaggcggag	gaggtaaaat	cgcctgcag	ccgaactatg	tcctgt <del>ctca</del>	aagtcc <del>tgac</del>	1080
aaagtgatct	taagaaattt	tgaaggtgt	attacgtcat	atccgg <del>aacc</del>	agagaattat	1140
atcccccaacc	aggcagacgc	ctat <del>ttt</del> gag	tccgtttcc	ctgaaa <del>accgc</del>	tgacaaaaag	1200
gagccgatcg	ggctgagtgc	cattttgct	gacaaagaag	tttcgt <del>ctac</del>	acctgaaaat	1260
gtagacagaa	tcaaacggcg	tgaggcatac	atcgcaaatc	cggag <del>c</del> atga	aacattaaaa	1320
gatcggcgtg	agaaaagggg	t <del>c</del> agctaaa	gaaaagaaaat	ttttggcgca	gcagaaaaaa	1380
cagaaaagaga	ctgaatgcgg	aggggattct	tcataa			1416

<210> 48  
 <211> 471  
 <212> PRT  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<400> 48

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Met Glu Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu  
1 5 10 15

Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln  
20 25 30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Ile Asp Pro Glu Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Ala Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Pro Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

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Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Cys Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
450 455 460

Glu Cys Gly Gly Asp Ser Ser  
 465 470

<210> 49  
 <211> 1416  
 <212> DNA  
 <213> Artificial Sequence  
  
 <220>  
 <223> Synthetic Construct  
  
 <400> 49  
 atgaaaaaca aatggtataa accgaaacgg cattggaagg agatcgagtt atggaaggac 60  
 gttccggaag agaaaatggaa cgattggctt tgacagctga cacacactgt aagaacgtta 120  
 gatgatttaa agaaaagtcat taatctgacc gaggatgaag aggaaggcgt ccgtatttct 180  
 accaaaaacga tccccttaaa tattacacct tactaggttt ctttaatgga ccccgacaaat 240  
 ccgagatgcc cggtacgcat gcagtctgtg ccactgtctg aagaaaatgca caaaaacaaaa 300  
 tacgataatgg aagacccgct tcatgaggat gaagattcac cggtacccgg tctgacacac 360  
 cgctatcccg accgtgtgct gtttcttgc acgaatcaat gttccgtgta ctgccgccac 420  
 tgcacacgccc ggcgcctttc cggacaaatc ggaatggcgcg tccccaaaaa acagcttgat 480  
 gctgcaattt cttatatccg gcaaacaccc gaaatccgcg attgttaat ttcaggcggt 540  
 gatgggctgc tcatcaacga ccaaattttttaa gaatataatt taaaagagct ggcgcacatt 600  
 ccgcacatctgg aagtcatccg catcgaaaca cgtgctcccg tcgtctttcc gcagcgcatt 660  
 accgatcatc tgtgcgagat attaaaaaaa tatcatccgg tctggctgaa cacccatttt 720  
 aacacaagca tcgaaaatgac agaagaatcc gttgaggcat gtgaaaagct ggtgaacgcg 780  
 ggagtgcgcg tcggaaatca ggctgtcgta ttagcaggta ttaatgattc ggttccaatt 840  
 atgaaaaagc tcatgcatttca cttggtaaaa atcagagtcc gtccttatttta tatttaccaa 900  
 tgtgatctgt cagaaggaat aaggcatttc cgtgctcctg tttccaaagg tttggagatc 960  
 attgaaggc tgagaggtca cacctcaggc aatgcggttc ccacctttgt cgttcacgca 1020  
 ccaggcggag gaggtaaaat cgccctgcag ccgaactatg tcctgtctca aagtcctgac 1080  
 aaagtgtatct taagaaaattt tgaagggtgt attacgtcat atccggaaacc agagaattat 1140  
 atccccaaatc aggccagacgc ctatttttag tccgtttcc ctgaaaaccgc tgacaaaaag 1200  
 gagccgatcg ggctgagtgc catttttgc gacaaagaag tttcgctac acctgaaaat 1260  
 gtagacagaa tcaaacggcg tgaggcatac atcgcaaattc cggagcatga aacattaaaa 1320  
 gatcggcgtg agaaaagagg tcagctaaa gaaaagaaaat ttttggcgca gcagaaaaaa 1380

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cagaaaagaga ctgaatgcgg aggggattct tcataa 1416

<210> 50  
<211> 71  
<212> PRT  
<213> Artificial Sequence

<220>  
<223> Synthetic Construct

&lt;400&gt; 50

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu  
1 5 10 15

Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln  
20 25 30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Asn Ile Thr Pro Tyr  
65 70

<210> 51  
<211> 399  
<212> PRT  
<213> Artificial Sequence

<220>  
<223> Synthetic Construct

&lt;400&gt; 51

Val Ser Leu Met Asp Pro Asp Asn Pro Arg Cys Pro Val Arg Met Gln  
1 5 10 15

Ser Val Pro Leu Ser Glu Glu Met His Lys Thr Lys Tyr Asp Met Glu  
20 25 30

Asp Pro Leu His Glu Asp Glu Asp Ser Pro Val Pro Gly Leu Thr His  
35 40 45

Arg Tyr Pro Asp Arg Val Leu Phe Leu Val Thr Asn Gln Cys Ser Val  
50 55 60

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Tyr Cys Arg His Cys Thr Arg Arg Arg Phe Ser Gly Gln Ile Gly Met  
65 70 75 80

Gly Val Pro Lys Lys Gln Leu Asp Ala Ala Ile Ala Tyr Ile Arg Glu  
85 90 95

Thr Pro Glu Ile Arg Asp Cys Leu Ile Ser Gly Gly Asp Gly Leu Leu  
100 105 110

Ile Asn Asp Gln Ile Leu Glu Tyr Ile Leu Lys Glu Leu Arg Ser Ile  
115 120 125

Pro His Leu Glu Val Ile Arg Ile Gly Thr Arg Ala Pro Val Val Phe  
130 135 140

Pro Gln Arg Ile Thr Asp His Leu Cys Glu Ile Leu Lys Lys Tyr His  
145 150 155 160

Pro Val Trp Leu Asn Thr His Phe Asn Thr Ser Ile Glu Met Thr Glu  
165 170 175

Glu Ser Val Glu Ala Cys Glu Lys Leu Val Asn Ala Gly Val Pro Val  
180 185 190

Gly Asn Gln Ala Val Val Leu Ala Gly Ile Asn Asp Ser Val Pro Ile  
195 200 205

Met Lys Lys Leu Met His Asp Leu Val Lys Ile Arg Val Arg Pro Tyr  
210 215 220

Tyr Ile Tyr Gln Cys Asp Leu Ser Glu Gly Ile Arg His Phe Arg Ala  
225 230 235 240

Pro Val Ser Lys Gly Leu Glu Ile Ile Glu Gly Leu Arg Gly His Thr  
245 250 255

Ser Gly Asn Ala Val Pro Thr Phe Val Val His Ala Pro Gly Gly  
260 265 270

Gly Lys Ile Ala Leu Gln Pro Asn Tyr Val Leu Ser Gln Ser Pro Asp  
275 280 285

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Lys Val Ile Leu Arg Asn Phe Glu Gly Val Ile Thr Ser Tyr Pro Glu  
 290 295 300

Pro Glu Asn Tyr Ile Pro Asn Gln Ala Asp Ala Tyr Phe Glu Ser Val  
 305 310 315 320

Phe Pro Glu Thr Ala Asp Lys Lys Glu Pro Ile Gly Leu Ser Ala Ile  
 325 330 335

Phe Ala Asp Lys Glu Val Ser Ser Thr Pro Glu Asn Val Asp Arg Ile  
 340 345 350

Lys Arg Arg Glu Ala Tyr Ile Ala Asn Pro Glu His Glu Thr Leu Lys  
 355 360 365

Asp Arg Arg Glu Lys Arg Gly Gln Leu Lys Glu Lys Lys Phe Leu Ala  
 370 375 380

Gln Gln Lys Lys Gln Lys Glu Thr Glu Cys Gly Gly Asp Ser Ser  
 385 390 395

<210> 52

<211> 1245

<212> DNA

<213> Artificial Sequence

<220>

<223> Synthetic Construct

<220>

<221> misc\_feature

<223> This parental sequence is a modification of the wild-type KAM of Clostridium stricklandii

<220>

<221> CDS

<222> (1)...(1245)

<400> 52

atg agt tta aag gat aag ttt ttt aca cat gta agc caa gaa gat tgg 48  
 Met Ser Leu Lys Asp Lys Phe Phe Thr His Val Ser Gln Glu Asp Trp  
 1 5 10 15

aat gat tgg aaa tgg caa gta aga aat cgt ata aag act gtt gaa gaa 96  
 Asn Asp Trp Lys Trp Gln Val Arg Asn Arg Ile Lys Thr Val Glu Glu  
 20 25 30

ctt aaa aaa tat att cca ctt act cca gaa gaa gaa ggg gta aaa 144  
 Leu Lys Tyr Ile Pro Leu Thr Pro Glu Glu Glu Gly Val Lys  
 35 40 45

cgc tgt ctt gat aca tta cgt atg gct att act cca tac tat cta tcg	192
Arg Cys Leu Asp Thr Leu Arg Met Ala Ile Thr Pro Tyr Tyr Leu Ser	
50 55 60	
cta att gat gta gaa aat cca aat gac cct gta aga aag caa gct gta	240
Leu Ile Asp Val Glu Asn Pro Asn Asp Pro Val Arg Lys Gln Ala Val	
65 70 75 80	
cct ctt tct tta gag ctg cat cgc gca gcg tct gat atg gaa gac cca	288
Pro Leu Ser Leu Glu Leu His Arg Ala Ala Ser Asp Met Glu Asp Pro	
85 90 95	
ctt cat gaa gat gga gat tct cca gtt cca gga ctt aca cat cgc tat	336
Leu His Glu Asp Gly Asp Ser Pro Val Pro Gly Leu Thr His Arg Tyr	
100 105 110	
cct gat cgc gtt ctt tta atg act gat caa tgt tca gta tac tgc	384
Pro Asp Arg Val Leu Leu Leu Met Thr Asp Gln Cys Ser Val Tyr Cys	
115 120 125	
cgc cac tgt act cgt aga cgc ttc gct ggt cga aca gat tct gct gtt	432
Arg His Cys Thr Arg Arg Phe Ala Gly Arg Thr Asp Ser Ala Val	
130 135 140	
gat acg aag caa ata gat gct gcg att gaa tat atc aaa aat act cca	480
Asp Thr Lys Gln Ile Asp Ala Ala Ile Glu Tyr Ile Lys Asn Thr Pro	
145 150 155 160	
caa gta aga gac gtt cta ctt tca gga gga gat gct cta tta atc tca	528
Gln Val Arg Asp Val Leu Leu Ser Gly Gly Asp Ala Leu Leu Ile Ser	
165 170 175	
gat gaa aag ctt gag tac aca atc aga aga ctt cgt gaa ata cca cac	576
Asp Glu Lys Leu Glu Tyr Thr Ile Arg Arg Leu Arg Glu Ile Pro His	
180 185 190	
gtt gag gtt att cgt att gga tca cgt gta cca gtt gta atg cca caa	624
Val Glu Val Ile Arg Ile Gly Ser Arg Val Pro Val Val Met Pro Gln	
195 200 205	
cgt att aca cca gaa cta gtt tct atg ctt aaa aag tat cat cca gta	672
Arg Ile Thr Pro Glu Leu Val Ser Met Leu Lys Lys Tyr His Pro Val	
210 215 220	
tgg tta aat aca cac ttc aac cat cct aat gaa att act gaa gag tct	720
Trp Leu Asn Thr His Phe Asn His Pro Asn Glu Ile Thr Glu Glu Ser	
225 230 235 240	
aaa cgt gca tgt gag tta ctt gct gat gca ggt att cct ctt gga aat	768
Lys Arg Ala Cys Glu Leu Leu Ala Asp Ala Gly Ile Pro Leu Gly Asn	
245 250 255	
caa agt gtg ctt ctt gca ggt gta aat gat tgc atg cac gtt atg aaa	816
Gln Ser Val Leu Leu Ala Gly Val Asn Asp Cys Met His Val Met Lys	
260 265 270	
aaa cta gta aat gac tta gtt aaa ata cgc gta cgt cct tac tat att	864

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Lys Leu Val Asn Asp Leu Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile				
275	280	285		
tat caa tgt gac ctt tca gtt gga att gag cac ttt cgc act cca gtt				912
Tyr Gln Cys Asp Leu Ser Val Gly Ile Glu His Phe Arg Thr Pro Val				
290	295	300		
gca aag gga ata gaa ata att gaa ggc tta aga gga cat act tca gga				960
Ala Lys Gly Ile Glu Ile Glu Gly Leu Arg Gly His Thr Ser Gly				
305	310	315	320	
tac tgc gtt cct aca ttt gtt gtg cat gca cct ggt ggt gga gga aaa				1008
Tyr Cys Val Pro Thr Phe Val Val His Ala Pro Gly Gly Gly Lys				
325	330	335		
act cca gtt atg cca aac tat gtt att tca caa aat cac aat aaa gtt				1056
Thr Pro Val Met Pro Asn Tyr Val Ile Ser Gln Asn His Asn Lys Val				
340	345	350		
att tta cgt aac ttt gaa ggt gta att aca act tac gat gag cct gat				1104
Ile Leu Arg Asn Phe Glu Gly Val Ile Thr Thr Tyr Asp Glu Pro Asp				
355	360	365		
cat tat act ttc cac tgt gac tgt gat gta tgc act gga aaa aca aat				1152
His Tyr Thr Phe His Cys Asp Cys Asp Val Cys Thr Gly Lys Thr Asn				
370	375	380		
gtt cat aag gtt gga gta gct gga ctt cta aat gga gag aca gcg aca				1200
Val His Lys Val Gly Val Ala Gly Leu Leu Asn Gly Glu Thr Ala Thr				
385	390	395	400	
ctt gaa cct gag ggt ttg gaa aga aaa caa aga gga cat cac taa				1245
Leu Glu Pro Glu Gly Leu Glu Arg Lys Gln Arg Gly His His				
405	410			

<210> 53  
 <211> 414  
 <212> PRT  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<400> 53

Met Ser Leu Lys Asp Lys Phe Phe Thr His Val Ser Gln Glu Asp Trp  
 1 5 10 15

Asn Asp Trp Lys Trp Gln Val Arg Asn Arg Ile Lys Thr Val Glu Glu  
 20 25 30

Leu Lys Lys Tyr Ile Pro Leu Thr Pro Glu Glu Glu Gly Val Lys  
 35 40 45

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Arg Cys Leu Asp Thr Leu Arg Met Ala Ile Thr Pro Tyr Tyr Leu Ser  
50 55 60

Leu Ile Asp Val Glu Asn Pro Asn Asp Pro Val Arg Lys Gln Ala Val  
65 70 75 80

Pro Leu Ser Leu Glu Leu His Arg Ala Ala Ser Asp Met Glu Asp Pro  
85 90 95

Leu His Glu Asp Gly Asp Ser Pro Val Pro Gly Leu Thr His Arg Tyr  
100 105 110

Pro Asp Arg Val Leu Leu Leu Met Thr Asp Gln Cys Ser Val Tyr Cys  
115 120 125

Arg His Cys Thr Arg Arg Arg Phe Ala Gly Arg Thr Asp Ser Ala Val  
130 135 140

Asp Thr Lys Gln Ile Asp Ala Ala Ile Glu Tyr Ile Lys Asn Thr Pro  
145 150 155 160

Gln Val Arg Asp Val Leu Leu Ser Gly Gly Asp Ala Leu Leu Ile Ser  
165 170 175

Asp Glu Lys Leu Glu Tyr Thr Ile Arg Arg Leu Arg Glu Ile Pro His  
180 185 190

Val Glu Val Ile Arg Ile Gly Ser Arg Val Pro Val Val Met Pro Gln  
195 200 205

Arg Ile Thr Pro Glu Leu Val Ser Met Leu Lys Lys Tyr His Pro Val  
210 215 220

Trp Leu Asn Thr His Phe Asn His Pro Asn Glu Ile Thr Glu Glu Ser  
225 230 235 240

Lys Arg Ala Cys Glu Leu Leu Ala Asp Ala Gly Ile Pro Leu Gly Asn  
245 250 255

Gln Ser Val Leu Leu Ala Gly Val Asn Asp Cys Met His Val Met Lys  
260 265 270

Lys Leu Val Asn Asp Leu Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile  
275 280 285

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Tyr Gln Cys Asp Leu Ser Val Gly Ile Glu His Phe Arg Thr Pro Val  
 290 295 300

Ala Lys Gly Ile Glu Ile Ile Glu Gly Leu Arg Gly His Thr Ser Gly  
 305 310 315 320

Tyr Cys Val Pro Thr Phe Val Val His Ala Pro Gly Gly Gly Lys  
 325 330 335

Thr Pro Val Met Pro Asn Tyr Val Ile Ser Gln Asn His Asn Lys Val  
 340 345 350

Ile Leu Arg Asn Phe Glu Gly Val Ile Thr Thr Tyr Asp Glu Pro Asp  
 355 360 365

His Tyr Thr Phe His Cys Asp Cys Asp Val Cys Thr Gly Lys Thr Asn  
 370 375 380

Val His Lys Val Gly Val Ala Gly Leu Leu Asn Gly Glu Thr Ala Thr  
 385 390 395 400

Leu Glu Pro Glu Gly Leu Glu Arg Lys Gln Arg Gly His His  
 405 410

<210> 54  
 <211> 1251  
 <212> DNA  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<220>  
 <221> CDS  
 <222> (1)..(125 1)

<400> 54  
 atg gca gaa agt cgt aga aag tat tat ttc cct gat gtc acc gat gag 48  
 Met Ala Glu Ser Arg Arg Lys Tyr Tyr Phe Pro Asp Val Thr Asp Glu  
 1 5 10 15

caa tgg tac gac tgg cat tgg cag gtc ctc aat cga att aag acg ctc 96  
 Gln Trp Tyr Asp Trp His Trp Gln Val Leu Asn Arg Ile Lys Thr Leu  
 20 25 30

gac cag ctg aaa aag tac gtt aca ctc acc gct gaa gaa gaa gag gga 144  
 Asp Gln Leu Lys Lys Tyr Val Thr Leu Thr Ala Glu Glu Glu Gly

35	40	45	
gta aaa gaa tcg ccc	aaa gta ctc cga	atg gct atc aca	cct tat tat
Val Lys Glu Ser Pro	Lys Val Leu Arg	Met Ala Ile Thr	Pro Tyr Tyr
50	55	60	
ttg agt ttg ata gac	ccc gag aat cct aat	tgt ccg att cgt	aaa caa
Leu Ser Leu Ile Asp	Pro Glu Asn Pro	Asn Cys Pro Ile	Arg Lys Gln
65	70	75	80
gcc att cct act caa	cag gaa ctg gta	cgt gct cct gaa	gat cag gta
Ala Ile Pro Thr Gln	Gln Glu Leu Val	Arg Ala Pro	Glu Asp Gln Val
85	90	95	
gac cca ctt agt gaa	gat gaa gat tcg	ccc gta ccc gga	ctg act cat
Asp Pro Leu Ser Glu	Asp Glu Asp Ser	Pro Val Pro	Gly Leu Thr His
100	105	110	
cgt tat ccg gat cgt	gta ttg ttc	ctt atc acg	gac aaa tgt tcg atg
Arg Tyr Pro Asp Arg	Val Leu Phe	Leu Ile Thr	Asp Lys Cys Ser Met
115	120	125	
tac tgt cgt cat tgt	act cgc cgt cgc	ttc gca gga	cag aaa gat gct
Tyr Cys Arg His Cys	Thr Arg Arg	Phe Ala Gly	Gln Lys Asp Ala
130	135	140	
tct tct cct tct gag	cgc atc gat cga	tgc att gac	tat ata gcc aat
Ser Ser Pro Ser Glu	Arg Ile Asp Arg	Cys Ile Asp	Tyr Ile Ala Asn
145	150	155	160
aca ccg aca gtc cgc	gat gtt ttg cta	tcg gga ggc	gat gcc ctc ctt
Thr Pro Thr Val Arg	Asp Val Leu Leu	Ser Gly Gly	Asp Ala Leu Leu
165	170	175	
gtc agc gac gaa cgc	ttg gaa tac ata	ttg aag cgt	ctg cgc gaa gta
Val Ser Asp Glu Arg	Leu Glu Tyr	Ile Leu Lys	Arg Leu Arg Glu Val
180	185	190	
cct cat gtg gag att	gtt cgt ata gga	agc cgt acg	ccg gta gtc ctc
Pro His Val Glu Ile	Val Arg Ile	Gly Ser Arg	Thr Pro Val Val Leu
195	200	205	
cct cag cgt ata acg	cct caa ttg gtg	gat atg ctc	aaa aaa tat cat
Pro Gln Arg Ile Thr	Pro Gln Leu Val	Asp Met Leu	Lys Lys Tyr His
210	215	220	
ccg gtg tgg ctg aac	act cac ttc aac	cac ccg aat	gaa gtt acc gaa
Pro Val Trp Leu Asn	Thr His Phe Asn	His Pro Asn	Glu Val Thr Glu
225	230	235	240
gaa gca gtg gag gct	tgt gaa aga atg	gcc aat gcc	ggt att ccg ttg
Glu Ala Val Glu Ala	Cys Glu Arg Met	Ala Asn Ala	Gly Ile Pro Leu
245	250	255	
ggt aac caa acg gtt	tta ttg cgt	gga atc aat	gat tgt aca cat gtg
Gly Asn Gln Thr Val	Leu Leu Arg	Gly Ile Asn Asp	Cys Thr His Val
260	265	270	

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atg aag aag ttg gta cat ttg ctg gta aag atg cgt gtg cgt cct tac Met Lys Arg Leu Val His Leu Leu Val Lys Met Arg Val Arg Pro Tyr 275 280 285	864
tat ata tat gta tgc gat ctt tcg ctt gga ata ggt cat ttc cgc acg Tyr Ile Tyr Val Cys Asp Leu Ser Leu Gly Ile Gly His Phe Arg Thr 290 295 300	912
ccg gta tct aaa gga atc gaa att atc gaa aat ttg cgc gga cac acc Pro Val Ser Lys Gly Ile Glu Ile Glu Asn Leu Arg Gly His Thr 305 310 315 320	960
tcg ggc tat gca gtt cct acc ttt gtg gta ggt gct ccg ggg ggt ggt Ser Gly Tyr Ala Val Pro Thr Phe Val Val Gly Ala Pro Gly Gly Gly 325 330 335	1008
ggt aag ata cct gta acg ccg aac tat gtt gta tct cag tcc cca cga Gly Lys Ile Pro Val Thr Pro Asn Tyr Val Val Ser Gln Ser Pro Arg 340 345 350	1056
cat gtg gtt ctt cgc aat tat gaa ggt gtt atc aca acc tat acg gag His Val Val Leu Arg Asn Tyr Glu Gly Val Ile Thr Thr Tyr Thr Glu 355 360 365	1104
ccg gag aat tat cat gag gag tgc gat tgt gag gac tgt cga gcc ggt Pro Glu Asn Tyr His Glu Glu Cys Asp Cys Glu Asp Cys Arg Ala Gly 370 375 380	1152
aag cat aaa gag ggt gta gct gca ctt tcc gga ggt cag cag ttg gct Lys His Lys Glu Gly Val Ala Ala Leu Ser Gly Gly Gln Gln Leu Ala 385 390 395 400	1200
atc gag oct tcc gac tta gct cgc aaa aaa cgc aag ttt gat aag aac Ile Glu Pro Ser Asp Leu Ala Arg Lys Lys Arg Lys Phe Asp Lys Asn 405 410 415	1248
taa	1251

<210> 55  
 <211> 416  
 <212> PRT  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<400> 55

Met Ala Glu Ser Arg Arg Lys Tyr Tyr Phe Pro Asp Val Thr Asp Glu  
 1 5 10 15

Gln Trp Tyr Asp Trp His Trp Gln Val Leu Asn Arg Ile Lys Thr Leu  
 20 25 30

Asp Gln Leu Lys Lys Tyr Val Thr Leu Thr Ala Glu Glu Glu Gly

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35

40

45

Val Lys Glu Ser Pro Lys Val Leu Arg Met Ala Ile Thr Pro Tyr Tyr  
50 55 60

Leu Ser Leu Ile Asp Pro Glu Asn Pro Asn Cys Pro Ile Arg Lys Gln  
65 70 75 80

Ala Ile Pro Thr Gln Gln Glu Leu Val Arg Ala Pro Glu Asp Gln Val  
85 90 95

Asp Pro Leu Ser Glu Asp Glu Asp Ser Pro Val Pro Gly Leu Thr His  
100 105 110

Arg Tyr Pro Asp Arg Val Leu Phe Leu Ile Thr Asp Lys Cys Ser Met  
115 120 125

Tyr Cys Arg His Cys Thr Arg Arg Arg Phe Ala Gly Gln Lys Asp Ala  
130 135 140

Ser Ser Pro Ser Glu Arg Ile Asp Arg Cys Ile Asp Tyr Ile Ala Asn  
145 150 155 160

Thr Pro Thr Val Arg Asp Val Leu Leu Ser Gly Gly Asp Ala Leu Leu  
165 170 175

Val Ser Asp Glu Arg Leu Glu Tyr Ile Leu Lys Arg Leu Arg Glu Val  
180 185 190

Pro His Val Glu Ile Val Arg Ile Gly Ser Arg Thr Pro Val Val Leu  
195 200 205

Pro Gln Arg Ile Thr Pro Gln Leu Val Asp Met Leu Lys Lys Tyr His  
210 215 220

Pro Val Trp Leu Asn Thr His Phe Asn His Pro Asn Glu Val Thr Glu  
225 230 235 240

Glu Ala Val Glu Ala Cys Glu Arg Met Ala Asn Ala Gly Ile Pro Leu  
245 250 255

Gly Asn Gln Thr Val Leu Leu Arg Gly Ile Asn Asp Cys Thr His Val  
260 265 270

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Met Lys Arg Leu Val His Leu Leu Val Lys Met Arg Val Arg Pro Tyr  
 275 280 285

Tyr Ile Tyr Val Cys Asp Leu Ser Leu Gly Ile Gly His Phe Arg Thr  
 290 295 300

Pro Val Ser Lys Gly Ile Glu Ile Glu Asn Leu Arg Gly His Thr  
 305 310 315 320

Ser Gly Tyr Ala Val Pro Thr Phe Val Val Gly Ala Pro Gly Gly  
 325 330 335

Gly Lys Ile Pro Val Thr Pro Asn Tyr Val Val Ser Gln Ser Pro Arg  
 340 345 350

His Val Val Leu Arg Asn Tyr Glu Gly Val Ile Thr Thr Tyr Thr Glu  
 355 360 365

Pro Glu Asn Tyr His Glu Glu Cys Asp Cys Glu Asp Cys Arg Ala Gly  
 370 375 380

Lys His Lys Glu Gly Val Ala Ala Leu Ser Gly Gly Gln Gln Leu Ala  
 385 390 395 400

Ile Glu Pro Ser Asp Leu Ala Arg Lys Lys Arg Lys Phe Asp Lys Asn  
 405 410 415

<210> 56  
 <211> 1278  
 <212> DNA  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<220>  
 <221> CDS  
 <222> (1)..(1278)

<400> 56  
 atg aat aca gtt aat act cgt aaa aaa ttt ttc cca aat gta act gat 48  
 Met Asn Thr Val Asn Thr Arg Lys Lys Phe Phe Pro Asn Val Thr Asp  
 1 5 10 15

gaa gaa tgg aat gat tgg aca tgg caa gta aaa aac cgc ctt aaa agt 96  
 Glu Glu Trp Asn Asp Trp Thr Trp Gln Val Lys Asn Arg Leu Lys Ser  
 20 25 30

gtt gaa gat tta gaa aaa tat gtt gat tta agt gaa gaa gaa aca gaa	144
Val Glu Asp Leu Glu Lys Tyr Val Asp Leu Ser Glu Glu Glu Thr Glu	
35 40 45	
ggg gtt gta cgc act ctt gaa act tta cgt atg gca atc act cca ttt	192
Gly Val Val Arg Thr Leu Glu Thr Leu Arg Met Ala Ile Thr Pro Phe	
50 55 60	
tac ttc tca ttg ata gat ttg aat agt gat cgc tgc cca ata cgt aag	240
Tyr Phe Ser Leu Ile Asp Leu Asn Ser Asp Arg Cys Pro Ile Arg Lys	
65 70 75 80	
caa gct ata cct act ata cga gaa ata cat caa tct gat gct gat atg	288
Gln Ala Ile Pro Thr Ile Arg Glu Ile His Gln Ser Asp Ala Asp Met	
85 90 95	
ttg gat cct cta cat gaa gat gaa gac tct cca gta cca gga tta act	336
Leu Asp Pro Leu His Glu Asp Glu Asp Ser Pro Val Pro Gly Leu Thr	
100 105 110	
cat cgc tat cca gat cgt gtt tta ctt cta ata aca gac atg tgt tct	384
His Arg Tyr Pro Asp Arg Val Leu Leu Ile Thr Asp Met Cys Ser	
115 120 125	
gta tac tgt cgc cac tgc act cgt cgc aga ttt gct ggg tca agt gat	432
Val Tyr Cys Arg His Cys Thr Arg Arg Arg Phe Ala Gly Ser Ser Asp	
130 135 140	
ggt gct atg cct atg gat aga att gac aaa gca ata gaa tat att gca	480
Gly Ala Met Pro Met Asp Arg Ile Asp Lys Ala Ile Glu Tyr Ile Ala	
145 150 155 160	
aaa act cca caa gta agg gat gta ttg tta tca gga gga gat gca ctt	528
Lys Thr Pro Gln Val Arg Asp Val Leu Leu Ser Gly Gly Asp Ala Leu	
165 170 175	
cta gtt tct aat aaa aaa tta gaa agc ata atc caa aaa cta cgc gca	576
Leu Val Ser Asn Lys Lys Leu Glu Ser Ile Ile Gln Lys Leu Arg Ala	
180 185 190	
ata cct cat gtt gaa ata atc aga ata gga agt cgt aca cca gtt gtt	624
Ile Pro His Val Glu Ile Ile Arg Ile Gly Ser Arg Thr Pro Val Val	
195 200 205	
tta cct caa aga att act cct gaa tta tgt aat atg tta aag aaa tat	672
Leu Pro Gln Arg Ile Thr Pro Glu Leu Cys Asn Met Leu Lys Lys Tyr	
210 215 220	
cat cca att tgg atg aat act cat ttt aac cac cct caa gaa gta acg	720
His Pro Ile Trp Met Asn Thr His Phe Asn His Pro Gln Glu Val Thr	
225 230 235 240	
cca gaa gct aaa aaa gct tgt gaa atg ttg gca gat gca gga gtt cca	768
Pro Glu Ala Lys Lys Ala Cys Glu Met Leu Ala Asp Ala Gly Val Pro	
245 250 255	
tta gga aat caa act gta cta tta aga gga ata aat gac agt gta cct	816

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Leu	Gly	Asn	Gln	Thr	Val	Leu	Leu	Arg	Gly	Ile	Asn	Asp	Ser	Val	Pro	
260								265						270		
gta atg aaa agg tta gta cat gat tta gta atg atg cgt gta cgc cct															864	
Val	Met	Lys	Arg	Leu	Val	His	Asp	Leu	Val	Met	Met	Arg	Val	Arg	Pro	
275								280					285			
tat tat att tac caa tgt gac tta tct atg gga ctc gaa cac ttc cgc															912	
Tyr	Tyr	Ile	Tyr	Gln	Cys	Asp	Leu	Ser	Met	Gly	Leu	Glu	His	Phe	Arg	
290					295					300						
aca cca gtt tct aaa ggt ata gaa att att gaa gga tta cgt gga cat															960	
Thr	Pro	Val	Ser	Lys	Gly	Ile	Glu	Ile	Glu	Gly	Leu	Arg	Gly	His		
305				310					315			320				
aca tct gga tat gca gta cca aca ttt gtt gtg cat gca cct ggt ggt															1008	
Thr	Ser	Gly	Tyr	Ala	Val	Pro	Thr	Phe	Val	Val	His	Ala	Pro	Gly	Gly	
325					330					335						
gga gga aaa act cca gta atg cct caa tat gta att tct caa tct cct															1056	
Gly	Gly	Lys	Thr	Pro	Val	Met	Pro	Gln	Tyr	Val	Ile	Ser	Gln	Ser	Pro	
340				345						350						
cat cgt gta gtt tta cgc aac ttt gaa gga gtt ata aca act tat aca															1104	
His	Arg	Val	Val	Leu	Arg	Asn	Phe	Glu	Gly	Val	Ile	Thr	Thr	Tyr	Thr	
355					360					365						
gaa cca gaa aat tat aca cat gaa cct tgt tat gat gaa gaa aaa ttt															1152	
Glu	Pro	Glu	Asn	Tyr	Thr	His	Glu	Pro	Cys	Tyr	Asp	Glu	Glu	Lys	Phe	
370				375						380						
gaa aaa atg tat gaa ata agt gga gtt tat atg cta gat gaa gga tta															1200	
Glu	Lys	Met	Tyr	Glu	Ile	Ser	Gly	Val	Tyr	Met	Leu	Asp	Glu	Gly	Leu	
385					390					395			400			
gaa atg tca cta gaa cct agc cac tta gca cgt cat gaa cgc aat aaa															1248	
Glu	Met	Ser	Leu	Glu	Pro	Ser	His	Leu	Ala	Arg	His	Glu	Arg	Asn	Lys	
405								410			415					
aag aga gca gaa gct gaa ggg aaa aaa taa															1278	
Lys	Arg	Ala	Glu	Ala	Glu	Gly	Lys	Lys								
420								425								

<210> 57  
 <211> 425  
 <212> PRT  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<400> 57

Met	Asn	Thr	Val	Asn	Thr	Arg	Lys	Lys	Phe	Phe	Pro	Asn	Val	Thr	Asp
1							5			10			15		

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Glu Glu Trp Asn Asp Trp Thr Trp Gln Val Lys Asn Arg Leu Lys Ser  
20 25 30

Val Glu Asp Leu Glu Lys Tyr Val Asp Leu Ser Glu Glu Glu Thr Glu  
35 40 45

Gly Val Val Arg Thr Leu Glu Thr Leu Arg Met Ala Ile Thr Pro Phe  
50 55 60

Tyr Phe Ser Leu Ile Asp Leu Asn Ser Asp Arg Cys Pro Ile Arg Lys  
65 70 75 80

Gln Ala Ile Pro Thr Ile Arg Glu Ile His Gln Ser Asp Ala Asp Met  
85 90 95

Leu Asp Pro Leu His Glu Asp Glu Asp Ser Pro Val Pro Gly Leu Thr  
100 105 110

His Arg Tyr Pro Asp Arg Val Leu Leu Leu Ile Thr Asp Met Cys Ser  
115 120 125

Val Tyr Cys Arg His Cys Thr Arg Arg Arg Phe Ala Gly Ser Ser Asp  
130 135 140

Gly Ala Met Pro Met Asp Arg Ile Asp Lys Ala Ile Glu Tyr Ile Ala  
145 150 155 160

Lys Thr Pro Gln Val Arg Asp Val Leu Leu Ser Gly Gly Asp Ala Leu  
165 170 175

Leu Val Ser Asn Lys Lys Leu Glu Ser Ile Ile Gln Lys Leu Arg Ala  
180 185 190

Ile Pro His Val Glu Ile Ile Arg Ile Gly Ser Arg Thr Pro Val Val  
195 200 205

Leu Pro Gln Arg Ile Thr Pro Glu Leu Cys Asn Met Leu Lys Lys Tyr  
210 215 220

His Pro Ile Trp Met Asn Thr His Phe Asn His Pro Gln Glu Val Thr  
225 230 235 240

Pro Glu Ala Lys Lys Ala Cys Glu Met Leu Ala Asp Ala Gly Val Pro  
245 250 255

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Leu Gly Asn Gln Thr Val Leu Leu Arg Gly Ile Asn Asp Ser Val Pro  
260 265 270

Val Met Lys Arg Leu Val His Asp Leu Val Met Met Arg Val Arg Pro  
275 280 285

Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser Met Gly Leu Glu His Phe Arg  
290 295 300

Thr Pro Val Ser Lys Gly Ile Glu Ile Glu Gly Leu Arg Gly His  
305 310 315 320

Thr Ser Gly Tyr Ala Val Pro Thr Phe Val Val His Ala Pro Gly Gly  
325 330 335

Gly Gly Lys Thr Pro Val Met Pro Gln Tyr Val Ile Ser Gln Ser Pro  
340 345 350

His Arg Val Val Leu Arg Asn Phe Glu Gly Val Ile Thr Thr Tyr Thr  
355 360 365

Glu Pro Glu Asn Tyr Thr His Glu Pro Cys Tyr Asp Glu Glu Lys Phe  
370 375 380

Glu Lys Met Tyr Glu Ile Ser Gly Val Tyr Met Leu Asp Glu Gly Leu  
385 390 395 400

Glu Met Ser Leu Glu Pro Ser His Leu Ala Arg His Glu Arg Asn Lys  
405 410 415

Lys Arg Ala Glu Ala Glu Gly Lys Lys  
420 425

<210> 58

<211> 1416

<212> DNA

<213> Artificial Sequence

<220>

<223> Synthetic Construct

<220>

<221> CDS

<222> (1)...(1416)

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<400> 58  
atg aaa aac aaa tgg tat aaa ccg aaa cg<sup>g</sup> cat tgg aag gag atc gag  
Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu  
1 5 10 15  
tta tgg aag gac gtt ccg gaa gag aaa tgg aac gat tgg ctt tgg cag 96  
Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln  
20 25 30  
ctg aca cac act gta aga acg tta gat gat tta aag aaa gtc att aat 144  
Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
35 40 45  
ctg acc gag gat gaa gag gaa ggc gtc cgt att tct acc aaa acg atc 192  
Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60  
ccc tta aat att aca cct tac tat gct tct tta atg gac ccc gac aat 240  
Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Asn  
65 70 75 80  
ccg aga tgc ccg gta cgc atg cag tct gtg ccg ctt tct gaa gaa atg 288  
Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95  
cac aaa aca aaa tac gat atg gaa gac ccg ctt cat gag gat gaa gat 336  
His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110  
tca ccg gta ccc ggt ctg aca cac cgc tat ccc gac cgt gtg ctg ttt 384  
Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125  
ctt gtc acg aat caa tgt tcc gtg tac tgc cgc cac tgc aca cgc cgg 432  
Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140  
cgc ttt tcc gga caa atc gga atg ggc gtc ccc aaa aaa cag ctt gat 480  
Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160  
gct gca att gct tat atc cgg gaa aca ccc gaa atc cgc gat tgt tta 528  
Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175  
att tca ggc ggt gat ggg ctg ctc atc aac gac caa att tta gaa tat 576  
Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190  
att tta aaa gag ctg cgc agc att ccg cat ctg gaa gtc atc cgc atc 624  
Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205  
gga aca cgt gct ccc gtc gtc ttt ccg cag cgc att acc gat cat ctg 672  
Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

tgc gag ata ttg aaa aaa tat cat ccg gtc tgg ctg aac acc cat ttt	720
Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe	
225 230 235 240	
aac aca agc atc gaa atg aca gaa gaa tcc gtt gag gca tgt gaa aag	768
Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys	
245 250 255	
ctg gtg aac gcg gga gtg ccg gtc gga aat cag gct gtc gta tta gca	816
Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala	
260 265 270	
ggt att aat gat tcg gtt cca att atg aaa aag ctc atg cat gac ttg	864
Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu	
275 280 285	
gta aaa atc aga gtc cgt cct tat att tac caa tgt gat ctg tca	912
Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser	
290 295 300	
gaa gga ata ggg cat ttc cgt gct cct gtt tcc aaa ggt ttg gag atc	960
Glu Gly Ile Gly His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile	
305 310 315 320	
att gaa ggg ctg aga ggt cat acc tca ggc tat gcg gtt cct acc ttt	1008
Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe	
325 330 335	
gtc gtt cac gca cca ggc gga gga ggt aaa atc gcc ctg cag ccg aac	1056
Val Val His Ala Pro Gly Gly Lys Ile Ala Leu Gln Pro Asn	
340 345 350	
tat gtc ctg tca caa agt cct gac aaa gtg atc tta aga aat ttt gaa	1104
Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu	
355 360 365	
ggt gtg att acg tca tat ccg gaa cca gag aat tat atc ccc aat cag	1152
Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln	
370 375 380	
gca gac gcc tat ttt gag tcc gtt ttc cct gaa acc gct gac aaa aag	1200
Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys	
385 390 395 400	
gag ccg atc ggg ctg agt gcc att ttt gct gac aaa gaa gtt tcg ttt	1248
Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Phe	
405 410 415	
aca cct gaa aat gta gac aga atc aaa cgg cgt gag gca tac atc gca	1296
Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala	
420 425 430	
aat ccg gag cat gaa aca tta aaa gat cgg cgt gag aaa aga gat cag	1344
Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Asp Gln	
435 440 445	
ctc aaa gaa aag aaa ttt ttg gcg cag cag aaa aaa cag aaa gag act	1392
Leu Lys Glu Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr	

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450

455

460

gaa tgc gga ggg gat tct tca taa  
 Glu Cys Gly Gly Asp Ser Ser  
 465 470

1416

<210> 59  
 <211> 471  
 <212> PRT  
 <213> Artificial Sequence

<220>  
 <223> Synthetic Construct

<400> 59

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu  
 1 5 10 15

Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln  
 20 25 30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
 35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
 50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Asn  
 65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
 85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
 100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
 115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
 130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
 145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu

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165

170

175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Gly His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

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Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Phe  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Asp Gln  
435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
450 455 460

Glu Cys Gly Gly Asp Ser Ser  
465 470

<210> 60  
<211> 471  
<212> PRT  
<213> lysine 2,3-aminomutase from *Bacillus subtilis*  
<400> 60

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu  
1 5 10 15

Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln  
20 25 30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Leu Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Arg Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

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Leu Val Thr Asn Gln Cys Ser Met Tyr Cys Arg Tyr Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Gly His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val Asp Ala Pro Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

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Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Phe  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Arg Arg Asp Gln  
435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
450 455 460

Glu Cys Gly Gly Asp Ser Ser  
465 470

<210> 61  
<211> 471  
<212> PRT  
<213> Artificial Sequence

<220>  
<223> Synthetic construct

<400> 61

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu  
1 5 10 15

Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln  
20 25 30 |

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

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Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Leu Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Met Tyr Cys Arg Tyr Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

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Glu Gly Ile Gly His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val Asp Ala Pro Gly Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Phe  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Asp Gln  
435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
450 455 460

Glu Cys Gly Gly Asp Ser Ser  
465 470

<210> 62

<211> 471

<212> PRT

<213> Artificial Sequence

<400> 62

Met Lys Asn Lys Trp Tyr Lys Pro Lys Arg His Trp Lys Glu Ile Glu  
1 5 10 15

Leu Trp Lys Asp Val Pro Glu Glu Lys Trp Asn Asp Trp Leu Trp Gln

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20

25

30

Leu Thr His Thr Val Arg Thr Leu Asp Asp Leu Lys Lys Val Ile Asn  
35 40 45

Leu Thr Glu Asp Glu Glu Gly Val Arg Ile Ser Thr Lys Thr Ile  
50 55 60

Pro Leu Asn Ile Thr Pro Tyr Tyr Ala Ser Leu Met Asp Pro Asp Asn  
65 70 75 80

Pro Arg Cys Pro Val Arg Met Gln Ser Val Pro Leu Ser Glu Glu Met  
85 90 95

His Lys Thr Lys Tyr Asp Met Glu Asp Pro Leu His Glu Asp Glu Asp  
100 105 110

Ser Pro Val Pro Gly Leu Thr His Arg Tyr Pro Asp Arg Val Leu Phe  
115 120 125

Leu Val Thr Asn Gln Cys Ser Val Tyr Cys Arg His Cys Thr Arg Arg  
130 135 140

Arg Phe Ser Gly Gln Ile Gly Met Gly Val Pro Lys Lys Gln Leu Asp  
145 150 155 160

Ala Ala Ile Ala Tyr Ile Arg Glu Thr Pro Glu Ile Arg Asp Cys Leu  
165 170 175

Ile Ser Gly Gly Asp Gly Leu Leu Ile Asn Asp Gln Ile Leu Glu Tyr  
180 185 190

Ile Leu Lys Glu Leu Arg Ser Ile Pro His Leu Glu Val Ile Arg Ile  
195 200 205

Gly Thr Arg Ala Pro Val Val Phe Pro Gln Arg Ile Thr Asp His Leu  
210 215 220

Cys Glu Ile Leu Lys Lys Tyr His Pro Val Trp Leu Asn Thr His Phe  
225 230 235 240

Asn Thr Ser Ile Glu Met Thr Glu Glu Ser Val Glu Ala Cys Glu Lys  
245 250 255

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Leu Val Asn Ala Gly Val Pro Val Gly Asn Gln Ala Val Val Leu Ala  
260 265 270

Gly Ile Asn Asp Ser Val Pro Ile Met Lys Lys Leu Met His Asp Leu  
275 280 285

Val Lys Ile Arg Val Arg Pro Tyr Tyr Ile Tyr Gln Cys Asp Leu Ser  
290 295 300

Glu Gly Ile Arg His Phe Arg Ala Pro Val Ser Lys Gly Leu Glu Ile  
305 310 315 320

Ile Glu Gly Leu Arg Gly His Thr Ser Gly Tyr Ala Val Pro Thr Phe  
325 330 335

Val Val His Ala Pro Gly Gly Lys Ile Ala Leu Gln Pro Asn  
340 345 350

Tyr Val Leu Ser Gln Ser Pro Asp Lys Val Ile Leu Arg Asn Phe Glu  
355 360 365

Gly Val Ile Thr Ser Tyr Pro Glu Pro Glu Asn Tyr Ile Pro Asn Gln  
370 375 380

Ala Asp Ala Tyr Phe Glu Ser Val Phe Pro Glu Thr Ala Asp Lys Lys  
385 390 395 400

Glu Pro Ile Gly Leu Ser Ala Ile Phe Ala Asp Lys Glu Val Ser Ser  
405 410 415

Thr Pro Glu Asn Val Asp Arg Ile Lys Arg Arg Glu Ala Tyr Ile Ala  
420 425 430

Asn Pro Glu His Glu Thr Leu Lys Asp Arg Arg Glu Lys Arg Gly Gln  
435 440 445

Leu Lys Glu Lys Lys Phe Leu Ala Gln Gln Lys Lys Gln Lys Glu Thr  
450 455 460

Glu Cys Gly Gly Asp Ser Ser  
465 470

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<211> 49  
<212> DNA  
<213> artificial sequence  
  
<220>  
<223> Bacillus specific primer

<220>  
<221> misc\_feature  
<223> Forward primer  
  
<400> 63  
ccagcctggc cataaggaga tatacatatg aaaaacaaat ggtataaac

49

<210> 64  
<211> 50  
<212> DNA  
<213> artificial sequence  
  
<220>  
<223> Bacillus specific primer

<220>  
<221> misc\_feature  
<223> Reverse primer  
  
<400> 64  
atggtgatgg tgatggtggc cagttggcc ttatgaagaa tccctccgc

50